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(71) Applicant (for all designated States except US): GENESENSE TECHNOLOGIES, INC. [CA/CA]; Sunnybrook HSC, Room S-115, 2075 Bayview Avenue, Toronto, Ontario M4N 3M5 (CA).

(72) Inventors; and

(75) Inventors/Applicants (for US only): WRIGHT, Jim, A. [CA/CA]; Apartment 902, 5418 Yonge Street, Toronto, Ontario M4N 6X4 (CA). YOUNG, Aiping, H. [CA/CA]; Apartment 508-88 Grandview Road, Toronto, Ontario M2N 6V4 (CA). DUGOURD, Dominique [CA/CA]; 2053 A Mt. Pleasant Road, Toronto, Ontario M4P 2M5 (CA).

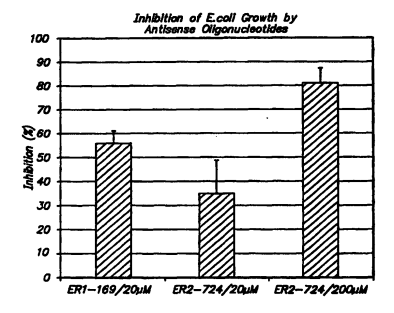
(74) Agent: DEETH WILLIAMS WALL; National Bank Building, Suite 400, 150 York Street, Toronto, Ontario M5H 3S5 (CA).

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(57) Abstract

The invention relates to antisense oligonucleotides which modulate the expression of the ribonucleotide reductase or the secA genes in microorganisms. This invention is als related to methods of using such oligonucleotides in inhibiting the growth of microorganisms. These antisense oligonucleotides are particularly useful in treating pathological conditions in mammals which are mediated by the growth of microorganisms.

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ANTISENSE OLIGONUCLEOTIDE SEQUENCES AS INHIBITORS OF MICROORGANISMS

BACKGROUND OF THE INVENTION

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Field of the Invention

This invention relates to antisense oligonucleotides which modulate the activity of the ribonucleotide reductase genes and the secA genes in microorganisms. This invention is also related to methods of using such compounds in inhibiting the growth of microorganisms.

These antisense oligonucleotides are particularly useful in treating pathological conditions in mammals which are mediated by the growth of microorganisms.

Accordingly, this invention also relates to pharmaceutical compositions comprising a pharmaceutically acceptable excipient and an effective amount of a compound of this invention.

These antisense oligonucleotides may also be used as anti-microbial agents for agricultural applications such as crop protection.

References

- The following publications, patent applications and patents are cited in this application as superscript numbers:
- 1. Nordlund and Eklund "Structure and function of the *Escherichia coli* ribonucleotide reductase protein R2", *J. Mol. Biol.* (1993) **232**:123-164;

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20

- 2. Carlson et al., "Primary structure of the *Escherichia coli* ribonucleoside diphosphate reductase operon", *PNAS* USA (1984) 81:4294-4297;
- 3. Nilsson et al., "Nucleotide sequence of the gene coding for the large subunit of ribonucleotide reductase of Escherichia coli Correction", Nucleic Acids Research (1988) 16:4174;
 - 4. P. Reichard, "The anaerobic ribonucleotide reductase from Escherichia coli", J. Biol. Chem. (1993) 268:8383-8386;

- 5. Nordlund et al., Nature (1990) 345:593-598;
- 6. der Blaauwen et al., "Inhibition of preprotein translocation and reversion of the membrane inserted state of secA by a carboxyl terminus binding Mab", *Biochemistry* (1997) 36:9159-9168;
- 7. McNicholas et al., "Dual regulation of Escherichia coli secA translation by distinct upstream elements", J. Mol. Biol. (1997) 265:128-141;
- 10 8. U.S. Patent No. 5,294,533;

5

15

- 9. Gasparro et al., "Photoactivatable antisense DNA: Suppression of ampicillin resistance in normally resistant Escherichia coli", Antisense Research and Development (1991) 1:117-140;
- 10. White et al., "Inhibition of the multiple antibiotic resistance (mar) operon in Escherichia coli by antisense DNA analogs", Antimicrobial Agents and Chemotherapy (1997) 41:2699-2704;
- 20 11. Nielsen et al., Science (1991) 354:1497;
 - 12. Good and Nielsen, "Inhibition of translation and bacterial growth by peptide nucleic acid targeted to ribosomal RNA", PNAS USA (1998) 95:2073-2076;
- 25 13. Buchardt, deceased, et al., U.S. Patent No. 5,766,855;
 - 14. Buchardt, deceased, et al., U.S. Patent No. 5,719,262;
 - 15. U.S. Patent No. 5,034,506;
 - 16. Altschul, et al., "Basic local alignment search tool", J. Mol. Biol. (1990) 215:403-10;
- 17. Devereux. et al., "A comprehensive set of sequence analysis programs for the VAX", Nucleic Acids Res. (1984) 12:387-395;
 - 18. Sambrook et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, New York (1989, 1992);
- 40 19. Ausubel et al., Current Protocols in Molecular Biology, John Wiley and Sons, Baltimore Maryland (1989);
 - 20. Chang et al., Somatic Gene Therapy, CRC Press, Ann Arbor MI (1995);

- 21. Vega et al., Gene Targeting, CRC Press, Ann Arbor MI (1995);
- 22. Vectors: A Survey of Molecular Cloning Vectors and Their Uses, Butterworths, Boston MA (1988)
- 23. U.S. Patent 5,023,252, issued June 11, 1991
- 24. Felgner et al., U.S. Patent No. 5,580,859.
- 10 25. U.S. Patent 5,011,472

5

- 26. Remington's Pharmaceutical Sciences, Mace Publishing Company, Philadelphia PA 17th ed. (1985);
- 15 27. Perbal, A Practical Guide to Molecular Cloning, John Wiley & Sons, New York (1988).
 - 28. PCR Protocols: A Guide To Methods And Applications, Academic Press, San Diego, CA (1990).
- 20 29. Dower, W.J., Nucleic Acids Res. (1988) 16:6127;
 - 30. Neuman et al., EMBO J. (1982) 1:841;
- 25 31. Taketo A., Biochim Biophys. Acta (1988) 949:318;
 - 32. Miller J.H. Experiments in Molecular Genetics, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y. (1972);
- 30 33. Horwitz J.P., J. Med. Chem. (1964) 7:574;
 - 34. Mann et al., Biochem. (1991) 30:1939;
 - 35. Olsvik, et al., Acta Pathol. Microbiol. Immunol. Scand. [B] (1982) 90:319;
- 36. Laemmli, U.K., Nature (1970) 227:680;
 - 37. Choy et al., Cancer Res. (1988) 48:2029;
- 40 38. Wright and Anazodo, Cancer J. (1988) 8:185-189;
 - 39. Chan et al., Biochemistry (1993) 32:12835-12840;
 - 40. Carpentier P.L., Microbiology 4th ed. W.B.Saunders Company (1977); and

41. Wright et al., Adv. Enzyme Regul. (1981) 19:105-127.

All of the above publications, patent applications and patents are herein incorporated by reference in their entirety to the same extent as if each individual publication, patent application or patent was specifically and individually indicated to be incorporated by reference in its entirety.

State of the Art

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Ribonucleotide reductase catalyzes the *de novo* production of deoxyribonucleotides. The enzyme reduces the four main ribonucleotides to the corresponding deoxyribonucleotides required for DNA synthesis and repair (Wright et al.⁴¹).

In mammalian and bacterial cells, *de novo* production of deoxyribonucleotides by ribonucleotide reductase is usually highly regulated on different levels in order to produce the correct amount of deoxyribonucleotides for DNA synthesis. In the DNA viruses, the metabolism of the host cell is directed towards production of viral DNA by virus encoded ribonucleotide reductases (Nordlund and Eklund¹).

Mammalian cells and many DNA viruses and prokaryotes, have a heterodimeric iron-containing ribonucleotide reductase enzyme of the $\alpha_2\beta_2$ type. For example, ribonucleotide reductase from E. coli is a multi-subunit $\alpha_2\beta_2$ enzyme where the two homo-dimeric proteins are denoted R1 and R2. The larger α_2 protein, R1, contains the binding sites for substrate and allosteric effectors and also the redox-active cysteine residues. Protein R1 has a molecular mass of 2 x 86,000 where each subunit contains 761 residues. The smaller β_2 protein, denoted R2, contains the dinuclear ferric center and a stable free tyrosyl radical necessary for the enzymatic activity. The R2 protein has a molecular mass of 2 x 43,500, where each subunit contains 375 amino acid residues (Nordlund and Eklund¹).

The nucleotide sequence of the *E. coli* K-12 DNA comprising the operon for the structural genes of the subunits of ribonucleotide reductase has been determined. The DNA sequence includes a total length of 8557 nucleotides. An open reading frame

between nucleotides 3506 and 5834 has been identified as the nrdA gene. An open reading frame between nucleotides 6012 and 7139 encoding a 375-amino acid polypeptide has been identified as the nrdB gene (Carlson et al.², and Nilsson et al.³). The sequences of the nrdA and nrdB genes for *E. coli* are shown in Figures 1 and 2.

In E. coli, the synthesis of ribonucleotide reductase is controlled at the level of transcription. The nrdA and nrdB genes direct the synthesis of a 3.2 kilobase polycistronic mRNA. Perturbations in DNA replication, either a shift up in growth conditions or an inhibition of DNA synthesis leads to increased synthesis of nrd mRNA (Carlson et al.²).

A separate anaerobic ribonucleotide reductase has also been identified from *E.coli*. The anaerobic *E. coli* reductase has a molecular mass of 145 kD and is a homodimer. The gene for the anaerobic reductase (nrdD) has been cloned and sequenced (P. Reichard⁴).

The ribonucleotide reductase R2 genomic or cDNA sequences are known for several other species such as bacteriophage T4, clam, mouse, Saccharomyces cerevisiae, vaccinia, herpes simplex virus types 1 and 2, varicella and Epstein-Barr virus (Nordlund et al.⁵). The sequence of the nrdE and nrdF which code for the ribonucleotide reductase genes of S. typhimurium are shown in Figure 3. The sequence of the ribonucleotide reductase gene of Lactococcus lactis is shown in Figure 4.

The secA gene of *E. coli* encodes for one component of a multi-component system for the secretion of proteins across the inner membrane of *E. coli* (der Blaauwen et al.⁶). The complete system consists of the SecB protein, a cytosolic chaperone, the SecA protein, the translocation ATPase and the heterotrimeric integral membrane SecY/SecE/SecG complex, which along with SecA serves as the preprotein channel. SecA protein plays a central role in the secretion process by binding the preprotein, secB protein, anionic phospholipids and SecY/SecE/SecG protein. These interactions allow SecA to recognize soluble preprotein and recruit it to translocation sites in the inner membrane. Once such protein translocation complexes have assembled; further steps require an ATP-driven cycle of insertion and de-insertion of

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secA protein with the inner membrane, where each cycle appears to be coupled to the translocation of a segment of the preprotein.

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SecA is the only component of the secretion apparatus that has been shown to be regulated. SecA is the second gene in the geneX-secA operon and its translation varies over a tenfold range depending on the status of protein secretion in the cell. During protein-export proficient conditions secA auto-represses its translation by binding to a site that overlaps the secA ribosome-binding site of genes-secA RNA. SecA protein can also dissociate a preformed 30 S-tRNA^{MET}-genes-secA RNA ternary complex in vitro. However, during a protein export block secA translation increases substantially although the mechanism responsible for this regulatory response has not been elucidated (McNicholas et al.⁷). The sequence of the secA gene of *E. coli* is shown in Figure 5.

The secA gene sequence has been identified for a number of other species including Mycobacterium bovis (Figure 6), Mycobacterium tuberculosis (Figure 7), Staphylococcus aureus (Figure 8), Staphylococcus carnosus (Figure 9), Bacillus subtilis, Bacillus firmus, Listeria monocytogenes, Mycobacterium smegmatis, Borrelia burgdorferi, P. sativum, S. griseus, and Synechoccus sp.

Antibiotics are important pharmaceuticals for the treatment of infectious diseases in a variety of animals including man. The tremendous utility and efficacy of antibiotics results from the interruption of bacterial (prokaryotic) cell growth with minimal damage or side effects to the eukaryotic host harboring the pathogenic organisms. In general, antibiotics destroy bacteria by interfering with the DNA replication, DNA to RNA transcription, translation (that is RNA to protein) or cell wall synthesis.

Although bacterial antibiotic resistance has been recognized since the advent of antimicrobial agents, the consequence of the emergence of resistant microorganisms, such resistance was historically controlled by the continued availability of effective alternative drugs. Now, drug resistance has emerged as a serious medical problem in the community, leading to increasing morbidity and mortality. The problem is worsened by the growing number of pathogens resistant to multiple, structurally

unrelated drugs. The situation has become so desperate that antibiotics once removed from use because of toxic effects may be prescribed in an attempt to deal with the otherwise untreatable drug resistant bacteria.

Antisense oligonucleotides have been used to decrease the expression of specific genes by inhibiting transcription or translation of the desired gene and thereby achieving a phenotypic effect based upon the expression of that gene (Wright and Anazado³⁸). For example, antisense RNA is important in plasmid DNA copy number control, in development of bacteriophage P22. Antisense RNA's have been used experimentally to specifically inhibit in vitro translation of mRNA coding specifically from Drosophila hsp23, to inhibit Rous sarcoma virus replication and to inhibit 3T3 cell proliferation when directed toward the oncogene c-fos. Furthermore, it is not necessary to use the entire antisense mRNA since a short antisense oligonucleotide can inhibit gene expression. This is seen in the inhibition of chloramphenicol acetyltransferase gene expression and in the inhibition of specific antiviral activity to vesicular stomatitus virus by inhibiting the N-protein initiation site. Antisense oligonucleotides directed to the macromolecular synthesis operon of bacteria, containing the rpsU gene, the rpoD gene and the dnaG gene have been used for the detection of bacteria. (U.S. Patent No. 5,294,5338). Furthermore, photoactivatable antisense DNA complementary to a segment of the β -lactamase gene has been used to suppress ampicillin resistance in normally resistant E. coli (Gasparro et al.9). Antisense DNA analogs have also been used to inhibit the multiple antibiotic resistant (mar) operon in *Escherichia coli* (White et al. 10).

Accordingly, there is a need to develop antisense oligonucleotides which will act to inhibit the growth of microorganisms.

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SUMMARY OF THE INVENTION

This invention is directed to antisense oligonucleotides which modulate the expression of the ribonucleotide reductase and secA genes in microorganisms and pharmaceutical compositions comprising such antisense oligonucleotides. This

invention is also related to methods of using such antisense oligonucleotides for inhibiting the growth of microorganisms.

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Accordingly, in one of its composition aspects, this invention is directed to an antisense oligonucleotide, which oligonucleotide is nuclease resistant and comprises from about 3 to about 50 nucleotides, which nucleotides are complementary to the ribonucleotide reductase gene or the secA gene of a microorganism. The antisense oligonucleotide may have one or more phosphorothioate internucleotide linkages.

In another of its composition aspects, this invention is directed to an antisense oligonucleotide comprising from about 3 to about 50 nucleotides which is capable of binding to the ribonucleotide reductase gene or the secA gene of a microorganism, wherein the oligonucleotide comprises all or part of a sequence selected from the group consisting of SEQ ID NO:22; SEQ ID NO:43; SEQ ID NO:62; SEQ ID NO:74; SEQ ID NO:75; SEQ ID NO:76; SEQ ID NO:143; SEQ ID NO:145; SEQ ID NO:152; SEQ ID NO:164; SEQ ID NO:176; SEQ ID NO:186; SEQ ID NO:188; SEQ ID NO:189; SEQ ID NO:191; SEQ ID NO:192; SEQ ID NO:195; SEQ ID NO:197; SEQ ID NO:206; SEQ ID NO:212; SEQ ID NO:220; SEQ ID NO:229; SEQ ID NO:235; SEQ ID NO:254; SEQ ID NO:261; SEQ ID NO:262; SEQ ID NO:263; SEQ ID NO:264; and SEQ ID NO:265.

In still another of its composition aspects, this invention is directed to a pharmaceutical composition comprising a pharmaceutically acceptable excipient and an effective amount of an antisense oligonucleotide, which oligonucleotide is nuclease resistant and comprises from about 3 to about 50 nucleotides, which nucleotides are complementary to the ribonucleotide reductase gene or the secA gene of a microorganism. The oligonucleotide may be modified, for example, the oligonucleotide may have one or more phosphorothioate internucleotide linkages.

In one of its method aspects, this invention is directed to a method for inhibiting the expression of the ribonucleotide reductase gene in a microorganism having a ribonucleotide reductase gene comprising, administering to said microorganism or to a cell infected with said microorganism an effective amount of an antisense oligonucleotide comprising from at least about 3 nucleotides which are complementary

to the ribonucleotide reductase gene of the microorganism under conditions such that the expression of the ribonucleotide reductase gene is inhibited.

In another of its method aspects, this invention is directed to a method for inhibiting the expression of the secA gene in a microorganism having a secA gene, comprising administering to said microorganism an effective amount of an antisense oligonucleotide comprising from at least about 3 nucleotides which are complementary to the secA gene of the microorganism under conditions such that expression of the secA gene is inhibited.

In one of its method aspects, this invention is directed to a method for inhibiting the growth of a microorganism encoding a ribonucleotide reductase gene or a secA gene, which method comprises administering to said microorganism or a cell infected with said microorganism an effective amount of an antisense oligonucleotide comprising from at least about 3 nucleotides which are complementary to either the ribonucleotide reductase gene or the secA gene of the microorganism under conditions such that the growth of the microorganism is inhibited. Preferably, the antisense oligonucleotide is selected from the group consisting of SEQ ID NO:22; SEQ ID NO:43; SEQ ID NO:62; SEQ ID NO:74; SEQ ID NO:75; SEQ ID NO:76; SEQ ID NO:143; SEQ ID NO:145; SEQ ID NO:152; SEQ ID NO:164; SEQ ID NO:176; SEQ ID NO:186; SEQ ID NO:188; SEQ ID NO:189; SEQ ID NO:191; SEQ ID NO:192; SEQ ID NO:195; SEQ ID NO:197; SEQ ID NO:206; SEQ ID NO:212; SEQ ID NO:220; SEQ ID NO:229; SEQ ID NO:235; SEQ ID NO:254; SEQ ID NO:261; SEQ ID NO:262; SEQ ID NO:263; SEQ ID NO:264; and SEQ ID NO:265.

In another of its method aspects, this invention is directed to a method for treating a mammalian pathologic condition mediated by a microorganism, which method comprises identifying a mammal having a pathologic condition mediated by a microorganism having a ribonucleotide reductase gene or a secA gene and administering to said mammal an effective amount of an antisense oligonucleotide comprising from at least about 3 nucleotides which are complementary to either the ribonucleotide reductase gene or the secA gene of the microorganism under conditions such that the growth of the microorganism is inhibited.

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BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is the sequence of the *E. coli* nrdA gene encoding the ribonucleotide reductase R1 subunit [SEQ ID NO:1].

Figure 2 is the sequence of the *E. coli* nrdB gene encoding the ribonucleotide reductase R2 subunit [SEQ ID NO:2]. The nrdB gene is encoded by nucleotides 7668 to 8798 of SEQ ID NO:2.

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Figure 3 is the sequence of the *S. typhimurium* nrdE and nrdF genes encoding the ribonucleotide reductase subunits [SEQ ID NO:3]. The nrdE gene is encoded by nucleotides 836 to 2980 and the nrdF gene is encoded by nucleotides 2991 to 3950 of SEQ ID NO:3.

Figure 4 is the sequence of the *Lactococcus lactis* nrdEF operon encoding ribonucleotide reductase [SEQ ID NO:4].

Figure 5 is the sequence of the E. coli secA gene [SEQ ID NO:5].

Figure 6 is the sequence of the Mycobacterium bovis secA gene [SEQ ID NO:6].

Figure 7 is the sequence of the *Mycobacterium tuberculosis* secA gene [SEQ ID NO:7].

Figure 8 is the sequence of the *Staphylococcus aureus* secA gene [SEQ ID NO:8].

Figure 9 is the sequence of the *Staphylococcus carnosus* secA gene [SEQ ID NO:9].

Figure 10 is the sequence of the bovine herpes virus ribonucleotide reductase small subunit gene [SEQ ID NO:10].

Figure 11 is the sequence of the Herpes simplex virus type 1 UL39 gene encoding ribonucleotide reductase 1 [SEQ ID NO:11].

Figure 12 is the sequence of the Herpes simplex type 2 ribonucleotide reductase gene [SEQ ID NO:12]. The ribonucleotide reductase gene is encoded by nucleotides 419 to 3853 of SEQ ID NO:12.

Figure 13 is the sequence of the equine herpes virus 4 ribonucleotide reductase large subunit and small subunit [SEQ ID NO:13]. The large subunit is encoded by

nucleotides 77 to 2446 and the small subunit by nucleotides 2485-3447 of SEQ ID NO:13.

Figure 14 is a photograph of a Western blot of a polyacrylamide gel of the cellular-protein-from E-coli-cells-carrying-a-plasmid-containing the mouse.

ribonucleotide reductase R2 gene after treatment with either $20\mu M$ or $200 \mu M$ of oligonucleotide AS-II-626-20.

Figure 15 is a graph of the inhibition of E. coli growth after treatment of E. coli cells with ribonculeotide reductase antisense oligonucleotides.

Figure 16 is a graph of the number of colony forming units/ml of E. coli cells after treatment with ribonucleotide reductase antisense oligonucleotides.

Figure 17 is a photograph of a Western blot of a polyacrylamide gel of cellular protein from *E. coli* cells after treatment with secA antisense oligonucleotides.

Figures 18a and 18b are graphs of the number of colony forming units/ml of E. coli cells after treatment with secA antisense oligonucleotides.

Figures 19a-g are graphs of growth curves of E. coli K12 after treatment with antisense oligonucleotides. Figure 19a shows the growth after treatment with 16 μ M or 80 μ M of antisense ES799 [SEQ ID NO:195]. Figure 19b shows the growth after treatment with 20 μ M of antisense ES1739 [SEQ ID NO:229]. Figure 19c shows the growth after treatment with 80 μ M of antisense ES851 [SEQ ID NO:197]. Figure 19d shows the growth after treatment with 80 μ M of antisense ES553 [SEQ ID NO:188]. Figure 19e shows the growth after treatment with 80 μ M of antisense ES646 [SEQ ID NO:191]. Figure 19f shows the growth after treatment with 80 μ M of antisense ES1845 [SEQ ID NO:235]. Figure 19g shows the growth after treatment with 80 μ M of antisense ES1845 [SEQ ID NO:235]. Figure 19g shows the growth after treatment with 80 μ M of antisense ES2537 [SEQ ID NO:254].

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DETAILED DESCRIPTION OF THE INVENTION

The present invention provides compounds that inhibit the growth of microbes by inhibiting the expression of a ribonucleotide reductase protein or the secA protein. Without being limited to any theory, the compounds inhibit the expression of the ribonucleotide reductase or the secA protein by inhibiting the transcription of the gene

or the translation of the mRNA to protein. Such compounds include antisense oligonucleotides.

Definitions:

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As used herein, the following terms have the following meanings:

The term "antisense oligonucleotide" as used herein means a nucleotide sequence that is complementary to the mRNA for the desired gene. Preferably, the antisense oligonucleotide is complementary to the mRNA for ribonucleotide reductase or secA.

The term "oligonucleotide" refers to an oligomer or polymer of nucleotide or nucleoside monomers consisting of naturally occurring bases, sugars, and inter-sugar (backbone) linkages. The term also includes modified or substituted oligomers comprising non-naturally occurring monomers or portions thereof, which function similarly. Such modified or substituted oligomers may be preferred over naturally occurring forms because of the properties such as enhanced cellular uptake, or increased stability in the presence of nucleases. The term also includes chimeric oligonucleotides which contain two or more chemically distinct regions. For example, chimeric oligonucleotides may contain at least one region of modified nucleotides that confer beneficial properties (e.g. increased nuclease resistance, increased uptake into cells) or two or more oligonucleotides of the invention may be joined to form a chimeric oligonucleotide.

The antisense oligonucleotides of the present invention may be ribonucleic or deoxyribonucleic acids and may contain naturally occurring or synthetic monomeric bases, including adenine, guanine, cytosine, thymine and uracil. The oligonucleotides may also contain modified bases such as xanthine, hypoxanthine, 2-aminoadenine, 6-methyl, 2-propyl and other alkyl adenines, 5-halo uracil, 5-halo cytosine, 6-aza uracil, 6-aza cytosine and 6-aza thymine, pseudo uracil, 4-thiouracil, 8-halo adenine, 8-aminoadenine, 8-thiol adenine, 8-thiolalkyl adenines, 8-hydroxyl adenine and other 8-substituted adenines, 8-halo guanines, 8-amino guanine, 8-thiol guanine, 8-thioalkyl guanines, 8-hydroxyl guanine and other 8-substituted guanines, other aza and deaza

uracils, thymidines, cytosines or guanines, 5-trifluoromethyl uracil and 5-trifluoro cytosine.

The antisense oligonucleotides of the invention may also comprise modified phosphorus oxygen heteroatoms in the phosphate backbone, short chain alkyl or cycloalkyl intersugar linkages or short chain heteroatom or heterocyclic intersugar linkages. For example, the antisense oligonucleotides may contain methyl phosphonates, phosphorothioates, phosphorodithioates, phosphotriesters, and morpholino oligomers. In one embodiment of the invention, the antisense oligonucleotides comprise phosphorothioate bonds linking between the four to six 3'-terminus nucleotides. In another embodiment, the phosphorothioate bonds link all the nucleotides. The antisense oligonucleotides may also have sugar mimetics.

The antisense oligonucleotides of the invention may also comprise nucleotide analogues wherein the structure of the nucleotide is fundamentally altered. An example of such an oligonucleotide analogue is a peptide nucleic acid (PNA) wherein the deoxyribose (or ribose) phosphate backbone in DNA (or RNA) is replaced with a polyamide backbone which is similar to that found in peptides (Nielsen et al.¹¹; Good and Nielsen¹²; Buchardt, deceased, et al.¹³, U.S. Patent No. 5,766,855; Buchardt, deceased, et al.¹⁴, U.S. Patent No. 5,719,262). PNA analogues have been shown to be resistant to degradation by enzymes and to have extended lives *in vivo* and *in vitro*. PNAs also bind more strongly to a complementary DNA sequence than to a naturally occurring nucleic acid molecule due to the lack of charge repulsion between the PNA strand and the DNA strand.

The oligonucleotides of the present invention may also include other nucleotides comprising polymer backbones, cyclic backbones, or acyclic backbones. For example, the nucleotides may comprise morpholino backbone structures (U.S. Patent No. 5,034,506¹⁵).

The oligonucleotides of the present invention are "nuclease resistant" when they have either been modified such that they are not susceptible to degradation by DNA and RNA nucleases or alternatively they have been placed in a delivery vehicle which in itself protects the oligonucleotide from DNA or RNA nucleases. Nuclease resistant

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oligonucleotides include, for example, methyl phosphonates, phosphorothioates, phosphorodithioates, phosphotriesters, and morpholino oligomers. Suitable delivery vehicles for conferring nuclease resistance include, for example liposomes.

The oligonucleotides of the present invention may also contain groups, such as groups for improving the pharmacokinetic properties of an oligonucleotides, or groups for improving the pharmacodynamic properties of an oligonucleotide. Preferably, the oligonucleotides do not contain reporter groups or labels, such as fluorescent dyes or radioactive labels.

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The antisense oligonucleotides may be complementary to the complete ribonucleotide reductase or secA gene including the introns. Preferably, the antisense oligonucleotides are complimentary to the mRNA region from the ribonucleotide reductase gene or the secA gene.

The antisense oligonucleotides may be selected from the sequence complementary to the ribonucletide reductase or secA genes such that the sequence exhibits the least likelihood of showing duplex formation, hair-pin formation, and homooligomer/sequence repeats but has a high to moderate potential to bind to the ribonucleotides reductase gene or the secA gene sequence and contains a GC clamp. These properties may be determined using the computer modeling program OLIGO Primer Analysis Software, Version 5.0 (distributed by National Biosciences, Inc., Plymouth, MN). This computer program allows the determination of a qualitative estimation of these five parameters.

Alternatively, the antisense oligonucleotides may also be selected on the basis that the sequence is highly conserved for either the ribonucleotide reductase or the secA genes between two or more microbial species. These properties may be determined using the BLASTN program (Altschul, et al. 16) of the University of Wisconsin Computer group (GCG) software (Devereux J. et al. 17) with the National Center for Biotechnology Information (NCBI) databases.

The antisense oligonucleotides generally comprise from at least about 3 nucleotides or nucleotide analogs, preferably from about 3 to about 50 nucleotides or

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nucleotide analogs, more preferably, from about 7 to about 35 nucleotides or nucleotide analogs, most preferably from about 15 to about 25 nucleotide or nucleotide analogs.

Preferably, the antisense oligonucleotides comprise from 3 to about 50 nucleotides or nucleotide analogs, more preferably from 20 to about 50 nucleotides or nucleotide analogs and further comprise all or part of the sequences set forth in Tables 1, 2, 3, and 4 (below). Preferably, the oligonucleotides complementary to the ribonucleotide reductase gene comprise SEQ ID NOS.: 14 to 157 as shown in Tables 1 and 2. Preferably, the antisense oligonucleotides complementary to the secA gene comprise the SEQ ID NOS.: 158 to 265 as shown in Tables 3 and 4.

Table 1

Antisense oligonucleotides that target the Escherichia coli K12 ribonucleotide reductase large subunit (R1)

L)	Milliona				large subulific (XXX)				
	Lego I	<u> </u>	Т		Sequence 5'-3'	Tm	(°C)	ΔG (kcal/mol)	
15	SEQ I		Name				1.1	-43.0	
	14		ER1-16	C	CGTCGCGCTTTGTCACCAG				
	-				TGTGCTACCGTCGCGCTTT	5	7.8	-42.0	
	15		ER1-24	1		1-	57.2	-40.2	
	16		ER1-33	Т	GATGCGCTCTGTGCTACCG	1		-	
20				+_	TTTGTCGAGATTGAT GCGCT		53.3	-38.7	
	17	7	ER1-44	1		- 1	51.7	-38.4	
	1	8	ER1-58	1	AGAACGCGATGGATTTTGTC				
	-			+	TGCCGCCCAATCCAGAACGC	7	64.6	-46.0	
	19	9	ER1-71	\perp		,+	57.7	-42.2	
		20	ER1-79	1	AGTCCTTCTGCCGCCCAATC				
				+	AAACTGAATGTGGGAGCGC	A	55.5	-39.8	
		21	ER1-128			_	55.5	-35.4	
. 25		22	ER1-169	9	ATAATGGTTTCGTGGATGT				
. <i>23</i>	-			\dashv	CGGCAGCCTTGATAATGGT	T	54.2	-40.6	
		23	ER1-18	U			51.4	-39.4	
		24 ER1-218		8	ATACTGATAATCCGGCGCA	11	31.4		
	-	24			TACGCAGGTGGAAGATCG	cc	57.3	-41.4	
		25	25 ER1-252		TACOCAGGT				

			Q ID	Na	me			 -			·			
			Vo: 26	ER1		-		uence 5			Tm	(°C)	ΔG (kca	l/mol)
		-		 				AGCG			64.	.4	4 -45.9	
		 	27 ER1-3			GCCC	CATCT	CGACC	CATTT	ГСА	54.	7	-39.7	,
			28 EI		330	TATO	GTAT	TTGCC	CATCT	CG	50.4	4	-38.1	
5		29		ER1-	123	CGGC	AGCAT	ΓAAGA	GAAGO	STC	51.6	5	-38.5	
5	5)	ER1-4	39	CCTTC	CCAGC	TGCT	TAACG	GC	56.4			
ı		`31		ER1-4	50	CCAGA							-41.9	
		32		ER1-4	79	ATAGA				- 1	51.5		-38.8	
		33		ER1-49						- 1	56.4		-41.8	
	3	34		ER1-50		GGAAC				- 1	53.9		-39.7	
10	 	35		ER1-51		GAATAT				- 1	48.5	T	-38.0	
	 	36				GCACGO				ı	52.2	1	-39.4	
•	-			R1-529		TCGAG	AACA	AGCAC	GCGG	С	60.8	\top	-43.3	-
	-			R1-543	T	TTCAC	GCGG	GTAGT	TCGAC	;	55.2	+	-40.5	\dashv
	-	38	E	R1-566	A	CGCTT	CACAT	TATTG	CAGGC	: -	52.2	+-	-38.7	_
		39	EF	R1-584	G	GAAAC	CGCG1	CGTA	AAAAC	+	3.9	┼		-
15	4	10	ER	1-592		TAAATG				+			-40.8	_
	4	1	ER	1-617	_	TGATT				+	2.7		-39.3	
	42	2	ER.	1-628		CACGC			_	64	1.0		-44.9	
	43	3	ERI	1-640					_	63	.8		-44.6	
	44			-667		AGTCG				64	.2		-45.8	1
20	45					GATCAC				52.	4	•	38.1	1
		-+-	ER1		GCT	GTCAC	CGCA	CTCGA	TCA	56.	9		39.1	
l	46		ER1-	689	GGA	ATCCA	GGCT	GTCAC	CGC	59.0			11.9	
							•							

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SEQ ID N:	Name	Sequence 5'→3'	Tm (°C)	ΔG (kcal/mol)
47	ER1-704	GGAGGTGGCGTTGATGGAAT	56.0	-40.6
48	ER1-7-16	AACAATCGCGCTGGAGGTGG	59.5	-42.7
49	ER1-778	CTACCCAGCGCACGAATACG	55.7	-40.9
50	ER1-817	ATGCAGCCGGTATGGAACGC	59.4	-43.1
51	ER1-829	TTGTAGAACGGAATGCAGCC	52.8	-38.8
52	ER1-846	CCGCTGTCTGGAAATGTTTG	53.1	-38.6
53	ER1-855	AGGATTTCACCGCTGTCTGG	54.0	-39.2
54	ER1-874	CGCACACCGCCCTGAGAGCA	63.9	-44.0
55	ER1-907	CACATCGGGTAGAACAGCGT	52.5	-38.1
56	ER1-925	CTTTCCACTTCCAGATGCCA	52.5	-38.1
57	ER1-964	TTGCCTTCCACACCACGGTT	57.5	-40.8
58	ER1-971	CACGCGGTTGCCTTCCACAC	60.8	-42.5
59	ER1-981	CCATATGACGCACGCGGTTG	59.4	-42.1
60	ER1-1034	TTCACCTTTCAGCAGACGGG	55.0	-39.6
61	ER1-1055	CGGGCTGAACAGGGTGATAT	53.8	-39.6
62	ER1-1059	CGGACGGCTGAACAGGGTG	62.1	-43.7
63	ER1-1061	GTCGGACGGGCTGAACAGGG	61.2	-43.4
64	ER1-1106	AAACTCTTCCTGATCGGCGA	53.8	-39.7
65	ER1-1148	GCGGATGCTGTCGTCTTTCT	54.3	-39.4
66	ER1-1155	GCTGCTTGCGGATGCTGTCG	61.3	-43.0
67	ER1-1166	GGCTTTCACACGCTGCTTGC	58.2	-41.4

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SEQ ID No:	Name	Sequence 5'→3'	Tm (°C)	ΔG (kcal/mol)
68	ER1-1173	GCTCAACGGCTTTCACACGC	58.0	-41.3
69	ER1-1212	GACCGGTAGACGCACGTTCC	56.7	-40.8
70	ER1-1255	GGGCTATGGGTATTGCAGTG	52.1	-38.7
71	ER1-1259	AAACGGGCTATGGGTATTGC	53.3	-40.7
72	ER1-1265	CGGATCAAACGGGCTATGGG	58.7	-43.4
73	ER1-1311	GGGCTATCTCCAGGCACAGG	55.9	-40.7
74	ER1-1315	GGCAGGGCTÄTCTCCAGGCA	58.7	-42.5
75	ER1-1320	TGGTCGGCAGGGCTATCTCC	58.6	-42.4
76	ER1-1326	GCGGTTTGGTCGGCAGGGCT	64.9	-47.0
77	ER1-1330	TTCAGCGGTTTGGTCGGCAG	60.5	-43.1
78	ER1-1336	ACGTCGTTCAGCGGTTTGGT	56.8	-40.9
79	ER1-1356	TTTCACCGTTCTCGTCGTTG	53.5	-38.5
80	ER1-1364	CAGCGCGATTTCACCGTTCT	57.5	-41.7
81	ER1-1370	CGTACACAGCGCGATTTCAC	54.2	-38.9
82	ER1-1379	AGCAGACAGCGTACACAGCG	54.0	-38.2
83	ER1-1388	CAGGTTGAAAGCAGACAGCG	53.4	-38.4
84	ER1-1397	AATTGCGCCCAGGTTGAAAG	56.5	-41.9
85	ER1-1407	CCAGGTTATTAATTGCGCCC	53.8	-41.3
86	ER1-1428	TTGCCAGCTCTTCCAGTTCA	53.3	-38.2
87	ER1-1438	ACCGCCAGAATTGCCAGCTC	58.8	-42.5
88	ER1-1451	GTCAAGTGCACGAACCGCCA	59.1	-41.0

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SEQ ID No:	Name	Sequence 5'-3'	Tm (°C)	ΔG (kcal/mol)
110	ER1-1974	CGTCGTGCAGGTGCTCGTAG	56.7	-39.9
111	ER1-1983	GCTCATAGGCGTCGTGCAGG	58.0	-41.4
112	ER1-1992	CCCACAGCAGCTCATAGGCG	58.0	-41.5
113	ER1-2000	CGGCATTTCCCACAGCAGCT	59.7	-42.8
114	ER1-2010	CATCGTTACCCGGCATTTCC	56.5	-41.9
115	ER1-2083	GGATCGTAGTTGGTGTTGGC	51.8	-39.9
116	ER1-2112	TCGGCACTTTTCCTGACGGG	59.5	-42.8
117	ER1-2145	AGGCGGTGAGCAGGTCTTTC	55.7	-40.5
118	ER1-2154	CGAATTTGTAGGCGGTGAGC	54.8	-40.5
119	ER1-2166	GTGTTTTGACCCCGAATTTG	51.9	-38.6
120	ER1-2211	CGTCTTGTGCGTCTTCAGCG	56.8	-40.0
121	ER1-2262	TCTTACATGCGCCGCTTTCG	58.6	-42.8

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Table 2
Antisense oligonucleotides that target the Escherichia coli K12 ribonucleotide reductase small subunit (R2)

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SEQ ID No:	Name	Sequence 5'→3'	Tm (°C)	ΔG (kcal/mol)
122	ER2-50	CGGCTGACCAAAGAACATCG	55.5	-40.0
123	ER2-60	CCACGTTGACCGGCTGACCA	61.2	-42.2
124	ER2-67	TAGCGAGCCACGTTGACCGG	60.6	-43.2
125	ER2-134	CGGACGCCAGAAGAAAGAGA	54.4	-39.8
126	ER2-144	CAACTTCTTCCGGACGCCAG	57.0	-41.3

	SEQ ID No:	Name	Sequence 5'-3'	Tm (°C)	ΔG (kcal/mol)
	127	ER2-168	AATCTATACGGTCGCGGGAG	53.4	-40.5
	128	ER2-198	TGTGTTTTTCGTGCTCCGGC	58.3	-41.6
	129	ER2-273	GCAATAGCGCCACGTTCGGG	62.1	-45.2
	130	ER2-284	AGAAATAAGCGGCAATAGCG	51.8	-40.3
5	131	ER2-290	CGGAATAGAAATAAGCGGCA	52.4	-40.3
	132	ER2-307	ACCCAGGTTTCCAGTTCCGG	57.4	-42.0
	133	ER2-350	ATAGGAACGGGAATGAATCG	50.7	-38.8
	134	ER2-441	TCCCTTCCGCACGTTTCTGG	59.5	-42.8
•	135	ER2-498	CGCCCAGCAGATGCCAGTAG	58.0	-41.5
10	136	ER2-505	GTACCTTCGCCCAGCAGATG	54.6	-39.7
	137	ER2-544	CGCAGGCTAACGGTCACAGT	55.2	-39.7
	138	ER2-557	TTTCTTCAGCTCGCGCAGGC	60.2	-43.4
	139	ER2-640	GCAAATGCGAAGGAACAAGC	54.9	-40.4
	140	ER2-655	ATCAATTCGCGTTCTGCAAA	53.4	-39.3
15	141	ER2-680	GCGAATAATTTTGGCGTTGC	. 54.9	-41.6
	142	ER2-692	GCGGGCAATCAGGCGAATAA	59.5	-44.0
	143	ER2-704	CAGGCTTCGTCGCGGCAA	66.8	-47.8
	144	ER2-714	CGGTCAGGTGCAGGGCTTCG	62.3	-44.0
	145	ER2-724	TGCTGGGTGCCGGTCAGGTG	63.6	-43.5
20	146	ER2-728	CATATGCTGGGTGCCGGTCA	58.8	-41.4
	147	ER2-778	GCAATTTCCGCCATCTCAGG	56.8	-41.5

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SEQ ID No:	Name	Sequence 5'→3'	Tm (°C)	ΔG (kcal/mol)
148	ER2-796	TCCTGCTTACACTCTTCGGC	52.1	-38.3
149	ER2-848	ATCCGCCCAGTCTTTCTCCT	54.2	-40.4
150	ER2-857	GAACAGATAATCCGCCCAGT	50.7	-38.1
151	ER2-976	GGGTTGGAGCGCGTCTGGAA	61.8	-44.0
152	ER2-983	CGGGATCGGGTTGGAGCGCG	68.1	-49.1
153	ER2-985	CACGGGATCGGGTTGGAGCG	64.0	-45.6
154	ER2-1045	CTGACTTCCACTTCCTGCGG	54.6	-39.9
155	ER2-1063	TGCCCGACCAGATAAGAACT	51.3	-38.2
156	ER2-1076	TTCCGAGTCAATCTGCCCGA	57.8	-41.2
157	ER2-1092	AATCGTCGGTGTCCACTTCC	53.6	-38.8

Table 3
Antisense Sequences that Target Escherichia coli SecA

SEQ ID No:	Name	Sequence 5 - 3'	Tm (°C)	ΔG kDa/mol
158	ES56	GACCACTTTGCGCATCCGGC	62.1	-44.2
159	ES62	GATGTTGACCACTTTGCGCA	54.3	-38.3
160	ES85	ATCTCCGGTTCCATGGCATT	55.5	-40.8
161	ES92	TTTTTCCATCTCCGGTTCCA	54.3	-40.1
162	ES116	CCCTTTCAGTTCTTCGTCGG	53.8	-39.8
163	ES124	GCGGTTTTCCCTTTCAGTTC	52.9	-39.9
164	ES129	ACTCTGCGGTTTTCCCTTTC	52.5	-39.6
165	ES153	CGCCTTTTTCCAGACGTGCA	58.4	-41.9
166	ES158	CACTTCGCCTTTTTCCAGAC	51.5	-38.4
167	ES165	TTTCCAGCACTTCGCCTTTT	54.1	-40.5

	SEQ ID No:	Name	Sequence 5 - 3'	Tm (°C)	ΔG kDa/mol
	168	ES170	CAGATTTTCCAGCACTTCGC	52.5	-38.6
	169	ES206	ACTTGCCTCACGTACCACGG	54.9	-39.5
	170	ES215	GACGCGCTTACTTGCCTCAC	55.0	-40.1
	171	ES230	GTGACGCATACCAAAGACGC	53.1	-38.5
5	172	ES264	TAAGAACCATACCGCCGAGT	51.5	-39.1
	173	ES286	ATTTCGGCGATGCAGCGTTC	59.7	-43.4
	174	ES303	TTCCTTCACCGGTACGCATT	54.5	-40.3
	175	ES307	GTTTTCCTTCACCGGTACG	51.4	-38.9
	176	ES320	CGTTGCGGTCAGGGTTTTTC	56.8	-41.6
10	177	ES336	TCAGGTAAGCAGGCAGCGTT	55.0	-40.2
	178	ES351	TACCGGTTAGTGCGTTCAGG	52.8	-39.2
	179	ES392	TTGCGCCAGGTAGTCGTTGA	56.5	-40.4
	180	ES398	GTCACGTTGCGCCAGGTAGT	55.0	-39.5
	181	ES418	AGCGGACGGTTGTTTTCGGC	60.8	-44.5
15	182	ES429	GGAATTCAAACAGCGGACGG	56.7	-41.5
	183	ES436	AGGCCAAGGAATTCAAACAG	51.0	-38.4
	184	ES448	ATACCGACAGTCAGGCCAAG	51.6	-38.0
	185	ES485	TTCGCGCTTTGCCGGTGCTG	65.8	-46.9
	186	ES531	AGCCGTATTCGTTGTTCGTA	50.1	-37.9
20	187	ES544	CGCAGGTAGTCAAAGCCGTA	53.1	-39.5
	188	ES553	ATGTTGTCGCGCAGGTAGTC	52.6	-38.1
	189	ES556	GCCATGTTGTCGCGCAGGTA	59.2	-41.7
	190	ES617	GTCCACTTCGTCCACCAGCG	57.7	-40.4
	191	ES646	GGTGTACGCGCTTCATCGAT	55.0	-40.0
25	192	ES647	CGGTGTACGCGCTTCATCGA	59.3	-42.1
	193	ES695	GCGTTTATACATTTCCGAGC	49.5	-38.4
	194	ES724	CGGATCAGGTGCGGAATAAT	53.9	-40.4

	SEQ ID N:	Name	Sequence 5 → 3'	Tm (°C)	ΔG kDa/mol
	195	ES799	TTCACCTGGCGAGATTTTTC	51.8	-38.6
	196	ES824	CAGCACCAGACCACGTTCGG	58.6	-40.7
	197	ES851	GCCCTCTTTCACCAGCAGTT	53.3	-39.1
	198	ES866	CCCTTCATCCATGATGCCCT	55.9	-40.6
5	199	ES889	TTGGCCGGAGAGTACAGAGA	52.2	-38.1
	200	ES898	AGCATGATGTTGGCCGGAGA	57.6	-40.9
	201	ES922	AGCGCCGCCGTTACGTGGTG	64.6	-46.5
	202	ES950	GTCACGGGTAAACAGCGCAT	54.9	-40.0
	203	ES1068	CACCTTCTTTCGCTTCCACA	52.8	-38.4
10	204	ES1097	CAGCGTTTGGTTTTCGTTCT	52.1	-38.9
	205	ES1109	GGTGATCGAAGCCAGCGTTT	56.5	-41.2
	206	ES1128	GACGGAAGTAGTTCTGGAAG	45.5	-35.0
	207	ES1147	CCCGCCAGTTTTTCATACAG	52.3	-39.2
	208	ES1152	TCATCCCCGCCAGTTTTTCA	57.5	-41.6
15	209	ES1218	GAACAACGACGGTATCCAGC	52.0	-38.2
	210	ES1328	GCCTTTCGCAGTACGTTCTT	51.4	-38.9
	211	ES1350	TAGTACCCACCAGCACCGGC	57.1	-41.4
	212	ES1398	CGGCTTTGGTCAGTTCGTTT	54.3	-40.1
	213	ES1410	TGTGCTTAATACCGGCTTTG	50.8	-38.6
20	214	ES1439	GTTGGCGTGGAATTTGGCGT	59.3	-43.0
	215	ES1462	GCCTGAGCAACAATCGCCGC	62.4	-44.5
	216	ES1515	CTGTACCACGACCCGCCATA	55.6	-40.3
	217	ES1518	TATCTGTACCACGACCCGCC	54.7	-40.0
	218	ES1545	CTGCCTGCCAGCTACCACCG	60.2	-42.9
25	219	ES1563	TTTCCAGCGCGGCAACTTCT	59.4	-43.4
	220	ES1581	TTTGCTCTGCGGTCGGATTT	57.0	-41.8
	221	ES1589	TTTTTCAATTTGCTCTGCGG	53.2	-39.8

SEQ ID No:	Name	Sequence 5 → 3'	Tm (°C)	ΔG kDa/m l
222	ES1624	ACCGCATCGTGACGTACCTG	55.7	-39.6
223	ES1629	CCAGTACCGCATCGTGACGT	<u>. 55.7</u>	39.6. mm
224	ES1633	GCTTCCAGTACCGCATCGTG	55.5	-40.0
225	ES1655	ACCGATGATATGCAGGCCAC	54.6	-39.6
226	ES1712	ACGACCAGAACGACCGCGCA	63.3	-44.1
227	ES1718	CCCCTGACGACCAGAACGAC	56.6	-40.1
228	ES1722	CATCCCCTGACGACCAGAA	56.9	-40.4
229	ES1739	GAAACGGGAAGAACCAGCAT	53.1	-39.5
230	ES1748	CGACAGGTAGAAACGGGAAG	51.4	-38.6
231	ES1781	GGAAGCAAAAATACGCATCA	50.6	-38.2
232	ES1785	GGTCGGAAGCAAAAATACGC	53.9	-40.9
233	ES1794	CGGATACTCGGTCGGAAGCA	57.3	-41.7
234	ES1814	ACCCAGTTTACGCATCATGC	52.5	-38.5
235	ES1845	ACGGGTGTTCAATGGCTTCG	57.1	-41.2
236	ES1861	ATCGCTTTAGTCACCCACGG	54.1	-40.0
237	ES1888	CTTTCAACTTTACGCTGGGC	51.9	-39.3
238	ES1892	ACGGCTTTCAACTTTACGCT	51.1	-39.2
239	ES2007	TGGTTTCGCTCACATCGCTG	57.0	-40.0
240	ES2054	GTAGGCATCAATGGTCGCTT	51.7	-38.5
241	ES2084	CCACATTTCTTCCAGCGACT	51.7	-38.0
242	ES2087	ATCCCACATTTCTTCCAGCG	53.9	-39.7
243	ES2191	TCACGCAGCGTCTCTTCATG	54.7	-38.2
244	ES2275	CCTTTCTCGAAGTGACGCAT	51.9	-38.2
. 245	ES2306	CCACAGGGAGTCAAGCGTTT	54.1	-39.3
246	ES2325	TCGCTGCCAGGTGCTCTTTC	57.7	-41.1
247	ES2330	GTCCATCGCTGCCAGGTGCT	59.7	-41.9
248	ES2339	ACGCAGATAGTCCATCGCTG	52.7	-38.4

SEQ ID No:	Name	Sequence 5 → 3'	Tm (°C)	ΔG kDa/mol
249	ES2381	CTTCGGATCTTTCTGTGCGT	51.9	-38.2
250	ES2395	CGTTTGTATTCCTGCTTCGG	52.5	-39.4
251	ES2422	ATCGCTGCAAACATGGAGAA	53.1	-38.5
252	ES2520	CCATACGACGCTGTTGTTCC	52.9	-38.5
253	ES2525	GGCTTCCATACGACGCTGTT	54.2	-40.0
254	ES2537	CGCTAAACGCTCGGCTTCCA	59.9	-44.1
255	ES2555	GCTAAGCTGCTGCATTTGCG	56.2	-41.3
256	ES2619	CTACTTTGCGCTCTCCGGTT	53.8	-40.4
257	ES2626	TTACGTCCTACTTTGCGCTC	50.0	-38.0
258	ES2646	AACCGCACGGGCAAGGATCG	63.6	-45.9
259	ES2651	ACCAGAACCGCACGGGCAAG	61.7	-44.0
260	ES2656	TTTTTACCAGAACCGCACGG	55.1	-41.0

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Table 4
Antisense Sequences that Target E. coli SecA based on Conserved Sequences

SEQ ID No:	Name	Sequence 5 → 3'	Tm (°C)	ΔG kDa/mol
261	ES386	CAGGTAGTCGTTGACGGTAA	47.7	-35.7
262	ES388	CAGGTAGTCGTTGACGGT	45.0	-32.9
263	ES1126	CGGAAGTAGTTCTGGAAGGT	47.6	-36.5
264	ES1702	CGACCGCGCAACTGGTTATC	57.8	-41.9
265	ES2644	CCGCACGGGCAAGGATCGTT	63.6	-45.9

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In Tables 1, 2, 3, and 4, the "Tm" is the melting temperature of an oligonucleotide duplex calculated according to the nearest-neighbor thermodynamic values. At this temperature 50% of nucleic acid molecules are in duplex and 50% are denatured. The " ΔG " is the free energy of the oligonucleotide, which is a measurement of an oligonucleotide duplex stability.

The following sequences have been determined to be conserved among species:

ES386 [SEQ ID NO:261] is conserved among Escherichia coli and

Mycobacterium tuberculosis;

ES388 [SEQ ID NO:262] is conserved among Escherichia coli; Mycobacterium tuberculosis; and Mycobacterium bovis;

ES553 [SEQ ID NO:188] is conserved among Escherichia coli, Mycobacterium tuberculosis, Mycobacterium bovis, Streptomyces coelicolor; and Streptomyces lividans;

ES556 [SEQ ID NO:189] is conserved among Escherichia coli, Mycobacterium tuberculosis, Mycobacterium bovis, Streptomyces coelicolor; and Streptomyces lividans; and Synechoccus sp.; and

ES646 [SEQ ID NO:191] is conserved among Escherichia coli and Staphylococcus carnosus;

ES1126 [SEQ ID NO:263] is conserved among *Escherichia coli* and *Rhodobacter capsulatus* SecA genes.

ES2644 [SEQ ID NO:265] is conserved among Escherichia coli SecA gene, MutA (A:T to C:G transversion), and tyrosine-specific transport protein (tyrP) gene.

The term "alkyl" refers to monovalent alkyl groups preferably having from 1 to 20 carbon atoms and more preferably 1 to 6 carbon atoms. This term is exemplified by groups such as methyl, ethyl, n-propyl, iso-propyl, n-butyl, iso-butyl, n-hexyl, and the like.

The term "aryl" refers to an unsaturated aromatic carbocyclic group of from 6 to 14 carbon atoms having a single ring (e.g., phenyl) or multiple condensed (fused) rings (e.g., naphthyl or anthryl). Preferred aryls include phenyl, naphthyl and the like.

The term "cycloalkyl" refers to cyclic alkyl groups of from 3 to 20 carbon atoms having a single cyclic ring or multiple condensed rings. Such cycloalkyl groups include, by way of example, single ring structures such as cyclopropyl, cyclobutyl, cyclopentyl, cyclooctyl, and the like, or multiple ring structures such as adamantanyl, and the like.

The term "halo" or "halogen" refers to fluoro, chloro, bromo and iodo and preferably is either fluoro or chloro.

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The term "thiol" refers to the group -SH.

As to any of the above groups which contain one or more substituents, it is understood, of course, that such groups do not contain any substitution or substitution patterns which are sterically impractical and/or synthetically non-feasible. In addition, the compounds of this invention include all stereochemical isomers arising from the substitution of these compounds.

The term "pharmaceutically acceptable salt" refers to salts which retain the biological effectiveness and properties of the antisense oligonucleotides of this invention and which are not biologically or otherwise undesirable. In many cases, the antisense oligonucleotides of this invention are capable of forming acid and/or base salts by virtue of the presence of amino and/or carboxyl groups or groups similar thereto.

Pharmaceutically acceptable base addition salts can be prepared from inorganic and organic bases. Salts derived from inorganic bases, include by way of example only, sodium, potassium, lithium, ammonium, calcium and magnesium salts. Salts derived from organic bases include, but are not limited to, salts of primary, secondary and tertiary amines, such as alkyl amines, dialkyl amines, trialkyl amines, substituted alkyl amines, di(substituted alkyl) amines, tri(substituted alkyl) amines, alkenyl amines, dialkenyl amines, trialkenyl amines, substituted alkenyl amines, di(substituted alkenyl) amines, tri(substituted alkenyl) amines, cycloalkyl amines, di(cycloalkyl) amines, tri(cycloalkyl) amines, substituted cycloalkyl amines, disubstituted cycloalkyl amine, trisubstituted cycloalkyl amines, cycloalkenyl amines, di(cycloalkenyl) amines, tri(cycloalkenyl) amines, substituted cycloalkenyl amines, disubstituted cycloalkenyl amine, trisubstituted cycloalkenyl amines, aryl amines, diaryl amines, triaryl amines, heteroaryl amines, diheteroaryl amines, triheteroaryl amines, heterocyclic amines, diheterocyclic amines, triheterocyclic amines, mixed di- and tri-amines where at least two of the substituents on the amine are different and are selected from the group consisting of alkyl, substituted alkyl, alkenyl, substituted alkenyl, cycloalkyl, substituted cycloalkyl, cycloalkenyl, substituted cycloalkenyl, aryl, heteroaryl,

heterocyclic, and the like. Also included are amines where the two or three substituents, together with the amino nitrogen, form a heterocyclic or heteroaryl group.

Examples of suitable amines include, by way of example only, isopropylamine, trimethyl amine, diethyl amine, tri(iso-propyl) amine, tri(n-propyl) amine, ethanolamine, 2-dimethylaminoethanol, tromethamine, lysine, arginine, histidine, caffeine, procaine, hydrabamine, choline, betaine, ethylenediamine, glucosamine, N-alkylglucamines, theobromine, purines, piperazine, piperidine, morpholine, N-ethylpiperidine, and the like. It should also be understood that other carboxylic acid derivatives would be useful in the practice of this invention, for example, carboxylic acid amides, including carboxamides, lower alkyl carboxamides, dialkyl carboxamides, and the like.

Pharmaceutically acceptable acid addition salts may be prepared from inorganic and organic acids. Salts derived from inorganic acids include hydrochloric acid, hydrobromic acid, sulfuric acid, nitric acid, phosphoric acid, and the like. Salts derived from organic acids include acetic acid, propionic acid, glycolic acid, pyruvic acid, oxalic acid, malic acid, malonic acid, succinic acid, maleic acid, fumaric acid, tartaric acid, citric acid, benzoic acid, cinnamic acid, mandelic acid, methanesulfonic acid, ethanesulfonic acid, p-toluene-sulfonic acid, salicylic acid, and the like.

The term "ribonucleotide reductase gene" or the "ribonucleoside diphosphate reductase gene" refers to any gene which encodes a protein that either reduces the four main ribonucleotides to the corresponding deoxyribonucleotides involved in DNA synthesis or encodes a subunit of a multimeric enzyme which reduces the four main ribonucleotides to the corresponding deoxyribonucleotides. Without being limiting, examples of ribonucleotide reductase genes from bacteria include the *E. coli* nrdA, nrdB and nrd D genes; the *S. typhimurium* nrdE and nrdF genes; and the *Lactococcus lactis* nrdEF gene. Examples of the ribonucleotide reductase genes from viruses include the herpes simplex type 1 and 2 ribonucleotide reductases and the bovine and equine herpes simplex ribonucleotide reductases.

The term "secA" refers to an oligonucleotide sequence which encodes a protein having similar properties as those expressed by the *E. coli* secA gene. Without being

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limiting, examples of secA genes from bacteria include the *Mycobacterium bovis* secA gene; the *Mycobacterium tuberculosis* secA gene, the *Staphylococcus aureus* secA gene and the *Staphylococcus carnosus* secA gene.

The term "microorganism" means a bacteria, fungi or virus having either a ribonucleotide reductase or secA gene. Specifically excluded from this definition is the malerial parasite, plasmodium.

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The term "bacteria" refers to any bacteria encoding either a ribonucleotide reductase gene or a secA gene, including Escherichi coli, Mycobacterium tuberculosis, Mycobacterium bovis, Mycobacterium smegmatis, Salmonella typhimurium, Thermoplasma acidophilum, Pyrococcus furiosus, Bacillus subtilis, Bacillus firmus, Lactococcus lactis, Staphylococcus aureus, Staphylococcus carnosus, Listeria monocytogenes, Borrelia burgdorferi, P. sativum, S. griseus, and Synechoccus sp.

The term "virus" refers to any virus having a ribonucleotide reductase gene.

Preferably the virus will be a DNA virus. Examples of suitable viruses include various herpes viruses (such as herpes simplex types 1 and 2, varicella-herpes zoster, cytomegalovirus and Epstein-Barr virus) and the various hepatitis viruses.

The term "complementary to" means that the antisense oligonucleotide sequence is capable of binding to the target sequence, ie the ribonucleotide reductase gene or the secA gene. Preferably the antisense oligonucleotide sequence has at least about 75% identity with the target sequence, preferably at least about 90% identity and most preferably at least about 95% identity with the target sequence allowing for gaps or mismatches of several bases. Identity can be determined, for example, by using the BLASTN program of the University of Wisconsin Computer Group (GCG) software.

The term "inhibiting growth" means a reduction in the growth of the bacteria or viruses of at least 25%, more preferably of at least 50% and most preferably of at least 75%. The reduction in growth can be determined for bacteria by a measuring the optical density of a liquid bacteria culture with a spectrophotometer or by counting the number of colony forming units/ml (CFU/ml) upon plating on culture plates. The reduction in growth can be determined for viruses by measuring the number of plaque forming units/ml upon plating on susceptible cells.

Preparation of the Antisense Oligonucleotides

The antisense oligonucleotides of the present invention may be prepared by conventional and well-known techniques. For example, the oligonucleotides may be prepared using solid-phase synthesis and in particular using commercially available equipment such as the equipment available from Applied Biosystems Canada Inc., Mississauga, Canada. The oligonucleotides may also be prepared by enzymatic digestion of the naturally occurring ribonucleotide reductase or secA gene by methods known in the art.

10 Isolation and Purification of the Antisense Oligonucleotides

Isolation and purification of the antisense oligonucleotides described herein can be effected, if desired, by any suitable separation or purification such as, for example, filtration, extraction, crystallization, column chromatography, thin-layer chromatography, thick-layer chromatography, preparative low or high-pressure liquid chromatography or a combination of these procedures. However, other equivalent separation or isolation procedures could, of course, also be used.

The invention contemplates a method of evaluating if an antisense oligonucleotide inhibits the growth of a microbe having a ribonucleotide reductase or secA gene. The method comprises selecting the microbe/microorganism having a ribonucleotide reductase or secA gene, administering the antisense oligonucleotide; and comparing the growth of the treated microbe with the growth of an untreated microorganism.

In order for the antisense oligonucleotide to effectively interrupt the expression of the ribonucleotide reductase or secA gene, the antisense oligonucleotide enters the microorganism's cell, in the case of fungal or bacterial cells or enter the mammalian cell having the virus target.

Although oligonucleotides are taken up by bacterial cells, some modification of the oligonucleotides may help facilitate or regulate said uptake. thus, a carier molecule, for example an amino acid, can be linked to the oligonucleotide. for example, bacteria have multiple transport systems for the recognition and uptake of

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molecules of leucine. The addition of this amino acid to the oligonucleotide may facilitate the uptake of the oligonucleotide in the bacteria and not substantially interfere with the activity of the antisense oligonucleotide in the bacterial cell.

Other methods are contemplated for facilitating the uptake of the antisense oligonucleotide into bacteria. For example, the addition of other amino acids or peptides or primary amines to the 3' or 5' termini of the antisense oligonucleotide may enable utilization of specific transport systems. Addition of lactose to the oligonucleotide by a covalent linkage may also be used to enable transport of the antisense oligonucleotide by lactose permease. Other sugar transport systems are also known to be functional in bacteria and can be utilized in this invention.

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With regard to inhibiting the expression of ribonucleotide reductase in DNA viruses, the antisense oligonucleotide is preferably introduced into the cell infected with the DNA virus. The antisense oligonucleotides may be delivered using vectors or liposomes.

An expression vector comprising the antisense oligonucleotide sequence may be constructed having regard to the sequence of the oligonucleotide and using procedures known in the art. The vectors may be selected from plasmids or benign viral vectors depending on the eukaryotic cell and the DNA virus. Phagemids are a specific example of beneficial vectors because they can be used either as plasmids or a bacteriophage vectors. Examples of other vectors include viruses such as bacteriophages, baculoviruses and retroviruses, DNA viruses, liposomes and other recombination vectors.

Vectors can be constructed by those skilled in the art to contain all the expression elements required to achieve the desired transcription of the antisense oligonucleotide sequences. Therefore, the invention provides vectors comprising a transcription control sequence operatively linked to a sequence which encodes an antisense oligonucleotide. Suitable transcription and translation elements may be derived from a variety of sources, including bacterial, fungal, viral, mammalian or insect genes. Selection of appropriate elements is dependent on the host cell chosen.

Reporter genes may be included in the vector. Suitable reporter genes include β -galactosidase (e.g. lacZ), chloramphenicol, acetyl-transferase, firefly luciferase, or an immunoglobulin or portion thereof. Transcription of the antisense oligonucleotide may be monitored by monitoring for the expression of the reporter gene.

The vectors can be introduced into cells or tissues by any one of a variety of known methods within the art. Such methods can be found generally described in Sambrook et al.¹⁸; Ausubel et al.¹⁹; Chang et al.²⁰; Vega et al.²¹; and Vectors: A Survey of Molecular Cloning Vectors and Their Uses²² and include, for example, stable or transient transfection, lipofection, electroporation and infection with recombinant viral vectors.

Introduction of nucleic acids by infection offers several advantages. Higher efficiency and specificity for tissue type can be obtained. Viruses typically infect and propagate in specific cell types. Thus, the virus' specificity may be used to target the vector to specific cell types *in vivo* or within a tissue or mixed culture of cells. Viral vectors can also be modified with specific receptors or ligands to alter target specificity through receptor mediated events.

Pharmaceutical Formulations

When employed as pharmaceuticals, the antisense oligonucleotides are usually administered in the form of pharmaceutical compositions. These compounds can be administered by a variety of routes including oral, rectal, transdermal, subcutaneous, intravenous, intramuscular, and intranasal. These compounds are effective as both injectable and oral compositions. Such compositions are prepared in a manner well known in the pharmaceutical art and comprise at least one active compound.

This invention also includes pharmaceutical compositions which contain, as the active ingredient, one or more of the antisense oligonucleotides associated with pharmaceutically acceptable carriers. In making the compositions of this invention, the active ingredient is usually mixed with an excipient, diluted by an excipient or enclosed within such a carrier which can be in the form of a capsule, sachet, paper or other container. When the excipient serves as a diluent, it can be a solid, semi-solid, or

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liquid material, which acts as a vehicle, carrier or medium for the active ingredient. Thus, the compositions can be in the form of tablets, pills, powders, lozenges, sachets, cachets, elixirs, suspensions, emulsions, solutions, syrups, aerosols (as a solid or in a liquid medium), ointments containing, for example, up to 10% by weight of the active compound, soft and hard gelatin capsules, suppositories, sterile injectable solutions, and sterile packaged powders.

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In preparing a formulation, it may be necessary to mill the active compound to provide the appropriate particle size prior to combining with the other ingredients. If the active compound is substantially insoluble, it ordinarily is milled to a particle size of less than 200 mesh. If the active compound is substantially water soluble, the particle size is normally adjusted by milling to provide a substantially uniform distribution in the formulation, e.g. about 40 mesh.

Some examples of suitable excipients include lactose, dextrose, sucrose, sorbitol, mannitol, starches, gum acacia, calcium phosphate, alginates, tragacanth, gelatin, calcium silicate, microcrystalline cellulose, polyvinylpyrrolidone, cellulose, sterile water, syrup, and methyl cellulose. The formulations can additionally include: lubricating agents such as talc, magnesium stearate, and mineral oil; wetting agents; emulsifying and suspending agents; preserving agents such as methyl- and propylhydroxy-benzoates; sweetening agents; and flavoring agents. The compositions of the invention can be formulated so as to provide quick, sustained or delayed release of the active ingredient after administration to the patient by employing procedures known in the art.

The compositions are preferably formulated in a unit dosage form, each dosage containing from about 5 to about 100 mg, more usually about 10 to about 30 mg, of the active ingredient. The term "unit dosage forms" refers to physically discrete units suitable as unitary dosages for human subjects and other mammals, each unit containing a predetermined quantity of active material calculated to produce the desired therapeutic effect, in association with a suitable pharmaceutical excipient. Preferably, the antisense oligonucleotide is employed at no more than about 20 weight percent of

the pharmaceutical composition, more preferably no more than about 15 weight percent, with the balance being pharmaceutically inert carrier(s).

The antisense oligonucleotide is effective over a wide dosage range and is generally administered in a pharmaceutically effective amount. It, will be understood, however, that the amount of the antisense oligonucleotide actually administered will be determined by a physician, in the light of the relevant circumstances, including the condition to be treated, the chosen route of administration, the actual compound administered, the age, weight, and response of the individual patient, the severity of the patient's symptoms, and the like.

For preparing solid compositions such as tablets, the principal active ingredient/antisense oligonucleotide is mixed with a pharmaceutical excipient to form a solid preformulation composition containing a homogeneous mixture of a compound of the present invention. When referring to these preformulation compositions as homogeneous, it is meant that the active ingredient is dispersed evenly throughout the composition so that the composition may be readily subdivided into equally effective unit dosage forms such as tablets, pills and capsules. This solid preformulation is then subdivided into unit dosage forms of the type described above containing from, for example, 0.1 to about 500 mg of the active ingredient of the present invention.

The tablets or pills of the present invention may be coated or otherwise compounded to provide a dosage form affording the advantage of prolonged action. For example, the tablet or pill can comprise an inner dosage and an outer dosage component, the latter being in the form of an envelope over the former. The two components can be separated by an enteric layer which serves to resist disintegration in the stomach and permit the inner component to pass intact into the duodenum or to be delayed in release. A variety of materials can be used for such enteric layers or coatings, such materials including a number of polymeric acids and mixtures of polymeric acids with such materials as shellac, cetyl alcohol, and cellulose acetate.

The liquid forms in which the novel compositions of the present invention may be incorporated for administration orally or by injection include aqueous solutions, suitably flavored syrups, aqueous or oil suspensions, and flavored emulsions with

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edible oils such as corn oil, cottonseed oil, sesame oil, coconut oil, or peanut oil, as well as elixirs and similar pharmaceutical vehicles.

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Compositions for inhalation or insufflation include solutions and suspensions in pharmaceutically acceptable, aqueous or organic solvents, or mixtures thereof, and powders. The liquid or solid compositions may contain suitable pharmaceutically acceptable excipients as described supra. Preferably the compositions are administered by the oral or nasal respiratory route for local or systemic effect. Compositions in preferably pharmaceutically acceptable solvents may be nebulized by use of inert gases. Nebulized solutions may be inhaled directly from the nebulizing device or the nebulizing device may be attached to a face mask tent, or intermittent positive pressure breathing machine. Solution, suspension, or powder compositions may be administered, preferably orally or nasally, from devices which deliver the formulation in an appropriate manner.

The following formulation examples illustrate representative pharmaceutical compositions of the present invention.

Formulation Example 1

Hard gelatin capsules containing the following ingredients are prepared:

20	Ingredient	Quantity (mg/capsule)
	Active Ingredient	30.0
	Starch	305.0
	Magnesium stearate	5.0
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The above ingredients are mixed and filled into hard gelatin capsules in 340 mg quantities.

Formulation Example 2

A tablet formula is prepared using the ingredients below:

		Quantity
	<u>Ingredient</u>	 (mg/tablet)
5	Active Ingredient	25.0
	Cellulose, microcrystalline	200.0
	Colloidal silicon dioxide	10.0
	Stearic acid	5.0

The components are blended and compressed to form tablets, each weighing

10 240 mg.

Formulation Example 3

A dry powder inhaler formulation is prepared containing the following components:

15	Ingredient	Weight %
	Active Ingredient	5
	Lactose	95

The active ingredient is mixed with the lactose and the mixture is added to a dry powder inhaling appliance.

Formulation Example 4

Tablets, each containing 30 mg of active ingredient, are prepared as follows:

25	Ingredient	Quantity (mg/tablet)
	Active Ingredient	30.0 mg
	Starch	45.0 mg
	Microcrystalline cellulose	35.0 mg
30	Polyvinylpyrrolidone	
	(as 10% solution in sterile water)	4.0 mg
	Sodium carboxymethyl starch	4.5 mg
	Magnesium stearate	0.5 mg
	Talc	_1.0 mg
35		<u>————</u>
	Total	120 mg

The active ingredient, starch and cellulose are passed through a No. 20 mesh U.S. sieve and mixed thoroughly. The solution of polyvinylpyrrolidone is mixed with the resultant powders, which are then passed through a 16 mesh U.S. sieve. The granules so produced are dried at 50° to 60°C and passed through a 16 mesh U.S. sieve. The sodium carboxymethyl starch, magnesium stearate, and talc, previously passed through a No. 30 mesh U.S. sieve, are then added to the granules which, after mixing, are compressed on a tablet machine to yield tablets each weighing 120 mg.

Formulation Example 5

Capsules, each containing 40 mg of medicament are made as follows:

	Ingredient	Quantity (mg/capsule)
15	Active Ingredient Starch Magnesium stearate Total	40.0 mg 109.0 mg <u>1.0 mg</u> 150.0 mg

The active ingredient, starch, and magnesium stearate are blended, passed through a No. 20 mesh U.S. sieve, and filled into hard gelatin capsules in 150 mg quantities.

Formulation Example 6

Suppositories, each containing 25 mg of active ingredient are made as follows:

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Ingredient	Amount
Active Ingredient Saturated fatty acid glycerides to	25 mg 2,000 mg

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The active ingredient is passed through a No. 60 mesh U.S. sieve and suspended in the saturated fatty acid glycerides previously melted using the minimum heat necessary. The mixture is then poured into a suppository mold of nominal 2.0 g capacity and allowed to cool.

Formulation Example 7

Suspensions, each containing 50 mg of medicament per 5.0 mL dose are made as follows:

5	Ingredient	Amount
	Active Ingredient	50.0 mg
	Xanthan gum	4.0 mg
	Sodium carboxymethyl cellulose (11%)	
	Microcrystalline cellulose (89%)	50.0 mg
10	Sucrose	1.75 g
	Sodium benzoate	10.0 mg
	Flavor and Color	q.v.
	Purified water to	5.0 mL

15 The active ingredient, sucrose and xanthan gum are blended, passed through a No. 10 mesh U.S. sieve, and then mixed with a previously made solution of the microcrystalline cellulose and sodium carboxymethyl cellulose in water. The sodium benzoate, flavor, and color are diluted with some of the water and added with stirring. Sufficient water is then added to produce the required volume.

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Formulation Example 8

	Ingredient	Quantity (mg/capsule)
25	Active Ingredient	15.0 mg
	Starch	407.0 mg
	Magnesium stearate	3.0 mg
	Total	425.0 mg

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The active ingredient, starch, and magnesium stearate are blended, passed through a No. 20 mesh U.S. sieve, and filled into hard gelatin capsules in 425.0 mg quantities.

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Formulation Example 9

A formulation may be prepared as follows:

	Ingredient		Quantity
5	Active Ingredient	•	5.0 mg
	Corn Oil		1.0 mL

Formulation Example 10

A topical formulation may be prepared as follows:

10	Ingredient	Quantity
	Active Ingredient	1-10 g
	Emulsifying Wax	30 g
15	Liquid Paraffin	20 g
	White Soft Paraffin	to 100 g

The white soft paraffin is heated until molten. The liquid paraffin and emulsifying wax are incorporated and stirred until dissolved. The active ingredient is added and stirring is continued until dispersed. The mixture is then cooled until solid.

Another preferred formulation employed in the methods of the present invention employs transdermal delivery devices ("patches"). Such transdermal patches may be used to provide continuous or discontinuous infusion of the antisense oligonucleotides of the present invention in controlled amounts. The construction and use of transdermal patches for the delivery of pharmaceutical agents is well known in the art. See, for example, U.S. Patent 5,023,252²³, herein incorporated by reference. Such patches may be constructed for continuous, pulsatile, or on demand delivery of pharmaceutical agents.

Another preferred method of delivery involves "shotgun" delivery of the naked antisense oligonucleotides across the dermal layer. The delivery of "naked" antisense oligonucleotides is well known in the art. See, for example, Felgner et al., U.S. Patent No. 5,580,859²⁴. It is contemplated that the antisense oligonucleotides may be packaged in a lipid vesicle before "shotgun" delivery of the antisense oligonucleotide.

Frequently, it will be desirable or necessary to introduce the pharmaceutical composition to the brain, either directly or indirectly. Direct techniques usually involve placement of a drug delivery catheter into the host's ventricular system to bypass the blood-brain barrier. One such implantable delivery system used for the transport of biological factors to specific anatomical regions of the body is described in U.S. Patent 5,011,472²⁵ which is herein incorporated by reference.

Indirect techniques, which are generally preferred, usually involve formulating the compositions to provide for drug latentiation by the conversion of hydrophilic drugs into lipid-soluble drugs. Latentiation is generally achieved through blocking of the hydroxy, carbonyl, sulfate, and primary amine groups present on the drug to render the drug more lipid soluble and amenable to transportation across the blood-brain barrier. Alternatively, the delivery of hydrophilic drugs may be enhanced by intra-arterial infusion of hypertonic solutions which can transiently open the blood-brain barrier.

Other suitable formulations for use in the present invention can be found in Remington's Pharmaceutical Sciences²⁶.

The antisense oligonucleotides or the pharmaceutical composition comprising the antisense oligonucleotides may be packaged into convenient kits providing the necessary materials packaged into suitable containers.

20 <u>Utility</u>

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The antisense oligonucleotides of the present invention may be used for a variety of purposes. They may be used to inhibit the expression of the ribonucleotide reductase gene in a microorganism, resulting in the inhibition of growth of that microorganism. They may be used to inhibit the expression of the secA gene in a microorganism, resulting in the inhibition of growth of that microorganism. The oligonucleotides may be used as hybridization probes to detect the presence of the ribonucleotide reductase gene or the secA gene in the microorganism. When so used the oligonucleotides may be labeled with a suitable detectable group (a radioisotope, a ligand, another member of a specific binding pair, for example, biotin). The oligonucleotides may also be used to determine the presence of a particular

microorganism in a biological sample. Finally, the oligonucleotides may be used as molecular wight markers.

In order to further illustrate the present invention and advantages thereof, the following specific examples are given but are not meant to limit the scope of the claims in any way.

EXAMPLES

In the examples below, all temperatures are in degrees Celsius (unless otherwise indicated) and all percentages are weight percentages (also unless otherwise indicated).

In the examples below, the following abbreviations have the following meanings. If an abbreviation is not defined, it has its generally accepted meaning:

 $\mu M = micromolar$

mM = millimolar

15 M = molar

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ml = milliliter

 $\mu l = microliter$

mg = milligram

 $\mu g = microgram$

20 IPTG = isopropyl- β -D-thiogalactoside

PAGE = polyacrylamide gel electrophoresis

PVDF = polyvinylidene difluoride

rpm = revolutions per minute

OD = optical density

25 CFU = colony forming units

 ΔG = free energy, a measurement of oligonucleotide duplex stability

kcal = kilocalories

General Methods in Molecular Biology:

Standard molecular biology techniques known in the art and not specifically described were generally followed as in Sambrook et al. 18; Ausubel et al. 19; and Perbal²⁷.

The antisense oligonucleotides in Tables 1, 2 and 3 were selected from the sequence complementary to the ribonucletide reductase or secA genes of *E. coli* such that the sequence exhibited the least likelihood of showing one or more of duplex formation, hair-pin formation, and homooligomer/sequence repeats but had a high to moderate potential to bind to the ribonucleotide reductase gene or the secA gene sequence. These properties were determined using the computer modeling program OLIGO Primer Analysis Software, Version 5.0 (distributed by National Biosciences, Inc., Plymouth, MN).

The antisense oligonucleotides in Table 4 were selected on the basis that the sequence is highly conserved for the secA genes between two or more microbial species. This property was determined using the BLASTN program (Altschul, et al. ¹⁶) of the University of Wisconsin Computer group (GCG) software (Devereux J. et al. ¹⁷) with the National Center for Biotechnology Information (NCBI) databases

Phosphorothioate oligonucleotides comprising the desired sequences were specially ordered either from Boston BioSystems, Bedford MA; Canadian Life Technologies, Burlington, Canada; Dalton Chemical Laboratories, Inc., North York, Canada; Hybridon, Inc., Milford Ma; Oligos Etc., or Oligos Therapeutics, Inc., Wilsonvill OR; or TriLink Bio Technologies, San Diego, CA. Antisense oligonucleotides may also be made by methods known in the art.

Polymerase chain reaction (PCR) was carried out generally as in PCR Protocols: A Guide To Methods And Applications²⁸.

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Example 1: Inhibition of mouse ribonucleotide reductase small subunit (R2) expression in Escherichia coli by antisense oligonucleotide AS-II-626-20

Competent BL21 (DE3) cells carrying a plasmid containing the mouse ribonucleotide reductase R2 gene were used. (Mann et al. 34) The antisense oligonucleotide, AS-II-626-20, GGCTAAATCGCTCCACCAAG [SEQ ID NO:266] is specifically complementary to the mouse ribonucleotide reductase R2 gene. Approximately 10^{10} bacteria/ml were electroporated using a Cell Porator (Gibco BRL, Burlington, Canada) in micro electro-chambers (0.4 cm between the electrodes) at a pulse of 2.4 kV, 4 k Ω with either 20 μ M or 200 μ M of antisense oligonucleotide AS-II-626-20, following methods described by the manufacturer (Dower W.J. 29 ; Neuman et; and Taketo, A. 31). Control populations were subjected to electroporation but without the antisense oligonucleotide AS-II-626-20.

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The bacterial cells were then transferred to Luria-Bertani broth (Miller J.H.³²) containing 50 μ g/ml of ampicillin and 0.4 mM of isopropyl β -D-thiogalactoside (IPTG) (expression inducer) (Horwitz J.P.³³) to grow at 30°C on a shaker at 250 rotations per minute (rpm) for 5 hours.

The cells were harvested by centrifugation and treated with 2 x sample loading buffer (100 mM Tris[hydroxymethyl'aminomethane, pH 6.8, 200 mM dithiothrietol, 4% sodium dodecyl sulfate, 20% glycerol and 0.015% bromophenol blue) and sonicated (Olsvik, et al.³⁵) for 15 seconds. The supernatants were resolved by polyacrylamide gel electrophoresis (PAGE) (Laemmli U.K.³⁶).

The ribonucleotide reductase R2 expression was detected by Western blot. The protein gel was blotted onto polyvinylidene difuoride (PVDF) protein sequencing membrane. (Choy et al.³⁷). The presence of the mouse ribonucleotide reductase was detected with a rabbit anti-mouse R2 subunit antibody (Chan et al.³⁹). The presence of the antibody bound to the ribonucleotide reducatase was detected using a second goat anti-rabbit immunoglobulin linked with horseradish peroxidase (Amersham Life Sciences, Oakville Canada).

The upper panel of Figure 14 is a photograph of the Western Blot results. The lower panel of Figure 14 is a photograph of the membrane stained with India ink to indicate the level of protein loaded in each lane.

It is clear that administration of either 20 \(^{\mu}M\) or 200 \(^{\mu}M\) AS-II-626-20 resulted, in a marked reduction of mouse ribonucleotide reductase gene expression in the E. coli cells.

Example 2: Inhibition of bacteria *Escherichia coli* K12 growth by antisense oligonucleotides ER1-169 and ER2-724 targeting *E. coli* ribonucleotide reductase large subunit (R1) and small subunit (R2)

E. coli cells were electroporated by the method set forth in Example 1 with ER1-169 [SEQ ID NO:22] or ER2-724 [SEQ ID NO:145] at the concentrations shown in Figure 15, while the control cells received oligonucleotide AS-II-626-20 [SEQ ID NO:] (targeting mouse ribonucleotide reductase small subunit).

The $E.\ coli$ cells were then transferred to fresh Luria-Bertani broth (Miller J.H. 32) to grow at 30°C on a shaker at 250 rpm for 3 hours. The flasks for the test and the control each contained the same number of bacteria per ml at the start of the experiment. The optical density at 590 nm (OD₅₉₀) of the cultures was measured at the start and at the end of the 3 hours. The inhibition of $E.\ coli$ growth was calculated by comparing the increase in OD₅₉₀ values at the start and the end of the 3 hours of the oligonucleotide-treated cultures to the increase of the control cultures at the start and at the end of the 3 hours. (Carpentier P.L. 40)

The results indicate that ER1-169 [SEQ ID NO:22] and ER2-724 [SEQ ID NO:145] inhibited the growth of *E. coli*.

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Example 3: Killing of Escherichia coli K12 by antisense oligonucleotides targeting the ribonucleotide reductase large subunit (R1) or the small subunit (R2)

E. coli cells (approximately 2 x 10^9 were incubated with 20 μ M of each of the phosphorothicate oligonucleotides set forth in Figure 12 on ice for 45 minutes. A

control without oligonucleotides was also incubated for each experiment. Cells were heat shocked by placing them in a 42°C bath for 45 seconds. (Sambrook J. et al. 18)

Luria-Bertani (LB) broth (Miller J.H.³²) was added and the samples were incubated at room temperature for 30 minutes. Dilutions of treated and untreated bacteria were incubated overnight at 37°C on culture plates containing LB medium, and the number of colonies was counted.

The number of killed bacteria was calculated by subtracting the surviving colony forming units (CFU/ml) of the oligonucleotide-treated bacteria from the CFU/ml of the control. Figure 16 shows the number of bacteria killed by treatment with the antisense sequences: ER1-640 [SEQ ID NO:43]; ER1-1059 [SEQ ID NO:62]; ER1-1320 [SEQ ID NO:75]; ER1-1315 [SEQ ID NO:74]; ER1-1326 [SEQ ID NO:76]; ER2-704 [SEQ ID NO:143] and ER2-983 [SEQ ID NO:152].

The results from Figure 16 show that antisense oligonucleotides complementary to either the R1 or R2 subunit of ribonucleotide reductase are effective as anti-bacterial agents.

Example 4: Inhibition of the secA protein expression in Escherichia coli following treatment with antisense phosphorothioate oligonucleotides

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E. coli cells were heat shock transformed by the method set forth in Example 3 above with the 80 μ M of each of the antisense phosphorothioate oligonucleotides set forth in Figure 17.

Luria-Bertani broth was then added to the treated *E. coli* cells and they were allowed to grow at 30°C on a shaker at 250 rpm for 3 hours.

Approximately the same quantity of treated and untreated bacteria, based on optical density, were washed in phosphate buffered saline, suspended in 2X Laemmli sample buffer (Laemmli U.K.³⁶), heated for 5 minutes at 95°C and subjected to SDS-PAGE (sodium dodecyl sulfate-polyacrylamide gel electrophoresis).

The gel was blotted onto polyvinylidene difluoride protein sequencing membrane by the methods set forth in Example 1. A rabbit polyclonal SecA antiserum (der Blaauwen et al.⁶) was used to detect the expression of the *E. coli* secA gene. The presence of bound rabbit antibody was detected using a goat anti-rabbit

5 immunoglobulin (Amersham Life Sciences, Oakville, Canada).

Figure 17 is a photograph of the Western Blot of *E. coli* cells treated with oligonucleotides ES799 [SEQ ID NO:195] (lane 1); ES1845 [SEQ ID NO:235] (lane 2); and the control (lane 3). When compared to the control, lane 3, the ES799 [SEQ ID NO:195] and ES1845 [SEQ ID NO:235] oligonucleotides clearly decreased the SecA protein levels in the treated *E. coli* cells. The top band in the Figure 17 represents SecA. Non-specific background bands appear below the SecA protein band.

It has been found that the antisense oligonucleotides are effective inhibitors of SecA expression in E. coli.

Example 5: Killing of Escherichia coli K12 by antisense secA oligonucleotides

E. coli cells were heat shock transformed by the method described in Example 3 above with either 100 μ M or 20 μ M of the antisense phosphorothioate oligonucleotides set forth in Figures 18a and 18b

Luria-Bertani (LB) broth (Miller J.H.³²) was added and the bacterial samples were incubated at room temperature for 30 minutes. Dilutions of treated and untreated bacteria were incubated overnight at 37°C on culture plates containing LB medium, and the number of colonies was counted.

The number of killed bacteria was calculated by subtracting the surviving colony forming units (CFU/ml) of the oligonucleotide-treated bacteria from the CFU/ml of the control. Figures 18a and 18b show the number of bacteria killed by treatment with the various antisense sequences. Accordingly, antisense oligonucleotides complementary to the secA gene act to inhibit the growth of *E. coli*.

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Example 6: Effect of antisense oligonucleotides on Escherichia coli K12 growth

E. coli cells were heat shock transformed by the method described in Example 3 with either 16 μ M, 20 μ M or 80 μ M of each of the antisense phosphorothioate oligonucleotides set forth in Figures 19a-g.

Equal numbers of the treated E. coli cells were then transferred to flasks containing fresh Luria-Bertani broth to grow at 30°C on a shaker at 250 rpm. The number of bacteria per flask was determined by the turbidity of the cultures at OD_{620} taken each hour (Carpentier P.L.⁴⁰).

Figures 19a-g show the rate of growth of the *E. coli* in each of the flasks after treatment with the various oligonucleotides. When growth curves of the treated and untreated cultures were statistically analyzed, the growth of the antisense treated cultures was found to be significantly inhibited when compared to the control cultures. The statistical p values are found in the Figures.

Claims:

1. An antisense oligonucleotide which is nuclease resistant and comprises from about 3 to about 50 nucleotides, which nucleotides are complementary to the ribonucleotide reductase gene or the seeA gene of a microorganism.

- 2. The oligonucleotide of Claim 1 comprising one or more phosphorothioate internucleotide linkages.
- An antisense oligonucleotide comprising from about 3 to about 50
 nucleotides which is capable of binding to the ribonucleotide reductase gene or the secA gene of a microorganism, wherein the oligonucleotide comprises all or part of a sequence selected from the group consisting of SEQ ID NO:22; SEQ ID NO:43; SEQ ID NO:62; SEQ ID NO:74; SEQ ID NO:75; SEQ ID NO:76; SEQ ID NO:143; SEQ ID NO:145; SEQ ID NO:152; SEQ ID NO:164; SEQ ID NO:176; SEQ ID NO:186;
 SEQ ID NO:188; SEQ ID NO:189; SEQ ID NO:191; SEQ ID NO:192; SEQ ID NO:195; SEQ ID NO:197; SEQ ID NO:206; SEQ ID NO:212; SEQ ID NO:220; SEQ ID NO:229; SEQ ID NO:235; SEQ ID NO:254; SEQ ID NO:261; SEQ ID NO:262; SEO ID NO:263; SEO ID NO:264; and SEQ ID NO:265.
- 4. A pharmaceutical composition comprising a pharmaceutically acceptable excipient and an effective amount of an oligonucleotide which is nuclease resistant and comprises from about 3 to about 50 nucleotides, which nucleotides are complementary to the ribonucleotide reductase gene or the secA gene of a microorganism.
- 25 5. The pharmaceutical composition comprising a pharmaceutically acceptable excipient and an effective amount of an oligonucleotide comprising from about 3 to about 50 nucleotides which is capable of binding to the ribonucleotide reductase gene or the secA gene of a microorganism, wherein the oligonucleotide comprises all or part of a sequence selected from the group consisting of SEQ ID NO:22; SEQ ID NO:43;
 30 SEQ ID NO:62; SEQ ID NO:74; SEQ ID NO:75; SEQ ID NO:76; SEQ ID NO:143;

SEQ ID NO:145; SEQ ID NO:152; SEQ ID NO:164; SEQ ID NO:176; SEQ ID NO:186; SEQ ID NO:188; SEQ ID NO:189; SEQ ID NO:191; SEQ ID NO:192; SEQ ID NO:195; SEQ ID NO:197; SEQ ID NO:206; SEQ ID NO:212; SEQ ID NO:220; SEQ ID NO:229; SEQ ID NO:235; SEQ ID NO:254; SEQ ID NO:261; SEQ ID NO:262; SEQ ID NO:263; SEQ ID NO:264; and SEQ ID NO:265.

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- 6. A method of inhibiting the expression of a ribonucleotide reductase gene in a microorganism having a ribonucleotide reductase gene, comprising administering to said microorganism or to a cell infected with said microorganism an effective amount of an antisense oligonucleotide comprising from at least about 3 nucleotides which are complementary to the ribonucleotide reductase gene of the microorganism under conditions such that the expression of the ribonucleotide reductase gene is inhibited.
- 7. The method according to Claim 6, wherein said microorganism is a bacterial cell.
 - 8. The method according to Claim 6, wherein said microorganism is a virus.
- 9. The method according to Claim 6 wherein the antisense oligonucleotide
 20 comprises a sequence selected from the group consisting of SEQ ID NO:22; SEQ ID
 NO:43; SEQ ID NO:62; SEQ ID NO:74; SEQ ID NO:75; SEQ ID NO:76; SEQ ID
 NO:143; SEQ ID NO:145; and SEQ ID NO:152.
- 10. A method of inhibiting the expression of the secA gene in a microorganism having a secA gene, comprising administering to said microorganism an effective amount of an antisense oligonucleotide comprising from at least about 3 nucleotides which are complementary to the secA gene of the microorganism under conditions such that the secA gene is inhibited.

11. The method according to Claim 10, wherein said microorganism is a bacterial cell.

12. The method according to Claim 11 wherein the antisense oligonucleotide comprises a sequence selected from the group consisting of SEQ ID NO:164; SEQ ID NO:176; SEQ ID NO:186; SEQ ID NO:188; SEQ ID NO:189; SEQ ID NO:191; SEQ ID NO:192; SEQ ID NO:195; SEQ ID NO:197; SEQ ID NO:206; SEQ ID NO:212; SEQ ID NO:220; SEQ ID NO:229; SEQ ID NO:235; SEQ ID NO:254; SEQ ID NO:265.

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- 13. A method of inhibiting the growth of a microorganism having a ribonucleotide reductase gene or a secA gene, which method comprises identifying the microorganism and administering to said microorganism an effective amount of an antisense oligonucleotide comprising from at least about 3 nucleotides which are complementary to either the ribonucleotide reductase gene or the secA gene of the microorganism under conditions whereby the growth of the microorganism is inhibited.
- 14. The method according to Claim 13, wherein said microorganism is a bacterial cell.

- 15. The method according to Claim 13, wherein said microorganism is a virus.
- 16. The method according to Claim 13 wherein the antisense oligonucleotide comprises a sequence selected from the group consisting of SEQ ID NO:22; SEQ ID NO:43; SEQ ID NO:62; SEQ ID NO:74; SEQ ID NO:75; SEQ ID NO:76; SEQ ID NO:143; SEQ ID NO:145; SEQ ID NO:152; SEQ ID NO:164; SEQ ID NO:176; SEQ ID NO:186; SEQ ID NO:188; SEQ ID NO:189; SEQ ID NO:191; SEQ ID NO:192; SEQ ID NO:195; SEQ ID NO:197; SEQ ID NO:206; SEQ ID NO:212; SEQ ID NO:220; SEQ ID NO:229; SEQ ID NO:235; SEQ ID NO:254; SEQ ID NO:261; SEQ ID NO:262; SEQ ID NO:263; SEQ ID NO:264; and SEQ ID NO:265.

17. A method for treating a mammalian pathologic condition mediated by microorganisms, which method comprises identifying a mammal having a pathologic condition mediated by microorganisms having a ribonucleotide reductase gene or a secA gene and administering to said mammal an effective amount of an antisense oligonucleotide comprising at least about 3 nucleotides which are complementary to either the ribonucleotide reductase gene or the secA gene of the microorganism under conditions such that the growth of the microorganism is inhibited.

gagatagece tgeegaecaa acegetgaae gaegteaaeg aegagaaegg tgaaategeg gtcagtctaa cctgtgcctg gctgttctcg tgaccactgc agtttgaacg tctgtatacc gaagatatca ccctgttcag cccgtccgac ggtgttgaaa acaaatcaac ctgcattccg gcgcggcggt cgccgggcgt ეედეენედიე cccgactcgt cgcgtgcttg caatatgtga agcgttttta cgacgcggtt acadaaccdc tcatctgctg cgaccgtgat tcagtatctc catecaedaa tgagccgcct gatttcccag cagagegeat caatetegae aageegttga ttcagaacgt ctggaagtgg aaagectget actacggggt tecagegega tigitaaata egiiteeeag egigeeggga teggeateaa gattegeggt ggtgaagegt teeataeegg atttecagae ageggtgaaa teetgetete agggeggtgt gecgaegeca ateatgteeg gegtgegtae tggattccat gagegeecag ttectttata ttetagttge atggacacct ttatcgatca gagatgggca aatacgataa ctggaaggca aatatctggt aaaaaagcct acggccagtt cgccggatta agacctctga ataacgitte gcgccagtgc cgtatctata gtgtggaagg caaccgcgtg cgtcatatgg gatcaggaag cagegtgtga gategagtge ggtgaeagee tecegtgatg gacggtatca gacggtagca gaaggactgc aatacccata gcccgtttga tccggccatc aaatatgaga aagacgacag cateegeaag ctgatgatge aggaacgtge gtetaeeggt tgtacgacge gttettegee atacccgtct gctgaaaggt tgttctaccc gatgtggcat aacgcgcctg atgacettet ettatgetge egttaageag gaaaatggtc agacctgatc ccacctgcgt gttcaagcag ttgggcggca teagttttat gacaaagcgc gtgaccggcg aaatctatga ttetegaact accegegtga tecacattta aaattteget gaagactaca cggaagaaga gccgcgcgc tggcgatctt gegetgtaeg aceaegtggt accattatca aggotgoogo gtcgagctgc gctcccacat gegttetgga atgaatcaga atctgctggt gtaccgggggc aaactgatgt ენაიიანნან ttctacaaac gacaaccgtg attcgtgcgc cagticaget gagatecate 081 841 901 781 721 481 541 601 661 301 361 241

FIG. 1A

ttacccgatc caacctgacg cctgctcdcc cgcatgtaag gatcaacttc gctggcgaaa cctgccgatc cgactgggaa tactetgata ggactacgag ttatctgcaa cactaaaaac actggaagag gccgcgcggt caactacqat acctggatga attatcagga tactgaaaga grattgaacc cccgtgacgg gegaaagegg cctctaatga cgctgcatta ccacgettte aggiggigco gtaacgatgg ctgccaacac deadedecaa gtattggtgt cgaaagggat gccactaacg gegetgetgg gcaattaata atgcagcagt tatcagaaca cgtacgctgg ctgctgaaag accacttacg getaatgage ctgcgtaact attttgcgcc tacteegaeg gasstgccgg gacgatggct cagtegatet caacctgggc cggtaaacgc tcagtattac aacactatat tacacttgac gatgggtegt qtttaacgaa gatetetaae gaaagacggt gtcaatccag ggataccatc aacgcacggt gctgctgtgg atttatcgat aaaagtgccg tggcggttcg cttcttcqca tecegteagg teggggtegg agtcaatcaa atctggtgcc tgtctgcttt aacgtggagc tggcgaacga cgtgcccgtg agaaagatet tcaaagcatc acacctatga tcatgcagaa tegaageeat ctggcaattc ctgtgtacgo pooboobboo ccgtccgaga tacqtcaqca cataaaacct gatacctata gcctacaaat deacaadaed qc t tac tacc gagcaaggcg gctctgcgtg cacctgcacg ctggtgggta ccgtcacgct atctga 441 921 501 681 861 2041 561 621 741 801 981 2101 2281

FIG. 11

actgtgaccg cgcgaattga cacctgaccg gcatatacca cageeggtea atteeggaae atcaaaataa aacgcactaa ctgatattqa ggcctgataa cacaggatge daaaadcadc gattaccagg acgetgetgg tcctatactc qtcaccaacg cttatccggc taacggtaaa atttqcagaa gatgagcgtt cgaageeetg teggtttgta gcgtgaacgc tegeateagg aaagctgatc cgaccgtata gettattet tcattcccqt tgacgatatc cgatgagetg catgtaagat attetttggt gaaatatcag cacactcatg ttgcccgcga cccacaccat daaadcddcd ggcctacggc cagaaacgat ctgttgtgtt ccagctatta atctctgcct attecttege gccggatgca cccaacadda aagaaccgat acatettega acgteteeg cactattacc cacaccaaca tcagcaacct attegeetga cacatettta ccgaacgtgg ggcgaaggta aaaaaactgt caatccagga cgatggctgc geggegtada egeettatee gatcagetea caaaaatatg tgggcgttct gaagggatet agetttgett cctgataaga gaagaagttg gcattagact aacgatccgt ggctccgggt tggaaggcaa cgccaaaatt gaaacgtgcg qcatctgctg cgagetgaag tttctacgtc gcacgaaaaa gggtcgtagc ggtcgaaacc cgtcgcatca gatttgtagg gccttatccg ctacgatcag ctggcgtccg taatatogtt gacgaaaaat agcagatca aagcgattcg tetetttett cgctgccgga attccattca atatcattcg ccagctactg ttagcctgcg ctggtgccgt acgtggctcg tggaaacctg gatgeeggat gacgcgccag ctacggctcg ggcgtaaaat cctttcaca 8101 8161 7381 7441 861 7981 8041 8281 7561 7621 7681 741 7801 921

FIG. 21

gtegeacaeg cgctggaatc cgatectgag atggeggada caacaddada attgactcgg aaagacattc ttggatctgc tctgataacg ccctgcgcat gcccgcgtta tcaggcagct ggtcgggcag cttctggcgg tgcggctcct ttggctggtg tggtctgaat ggcagtcggt cgaaggttac acaacacaaa acctgtttgt gttcgatgat acaccettee tccgtatgca ggatcaacac agetetgatg gttcttatct accagigica tatgctgaat ctgctgcgca gagtgctatg atcaccaata gtggaagtca agtaacttcc gccaggatga ttccgcgacg ccgatcccgt gtgtaagcag ggattatetg cgttgaatac gegetecaae tccgcaggaa cgacgatttg geggttgagt caactgctgt dcacccadca cactggcaca ccacaatgtg tgcaggttgc ttgccgaaga tctgccagta cgttccagac aagtggacac aagactgggc 8881 8461 8641 8701 8401 8521 8581 8761 8821

FIG. 2E

agggtgacga ccgctacccc tcctqctccq aactctccaa cgccgatcaa aagacgcatt tccatacaca aaaaatccg aggegatega gcaaacgtta gggcgcaagg atttcatcaa cgtattccgc taccgctttg agcatgaacg attacgaccg tecagaegtt catgattaac ttctccaaca ccgtcatacg aatgatgaac gaatttagge ggcgatgcct tgaacgcgat gtctcctgct teggegetge geggittate cactttcaac aaaatgotgg aacgetgaeg 6066606606 tggtetgget, teegeeegga taccacaccc aaddaccadc titgccagcc qtcctcgcgc ggettteget accttcgacg acgetggege getegtetae gcctgccacg tctggttgtg cegettttta tegeaattte cccctggctg aggagtaaat cadacdadda tggggaactg cccggtgatg 2662266666 aatgeggegt atgiccgaaa cqatgacgcc cacgetgaaa ggtggcgttg getttetggt ggcggtgaat caatctgcgc gatacactcc cctccggtaa aaccatggat teagttegae ttccgtgacg ccatgccagc aaccatacat gcatgagcgc gacaggtgat gtctggggct ccggcgtgat poobobobob ttctggatac cgatttgagc ttatqcaqc atagcacaaa gacategata tecqeeeqea aagggtatta ataccagtta ccgatgaaat aacadcadcd cgatcgggcg ttttactctc gggtgacaat ggatgcagcg odeddeadac caggtagacg ggcgttatcg taatgcagga caggccatat tegageaege gcggtgcgc attctgcgtt tttgaagatc acccaactga aattgeggea aatcagtctt aaccaactta tacgcaccgc tggcgatgtg tacgataaag decaeecaed ctggttcggg ggcgtcgcgt gegeattege tggaagttet aacatggagt atctggtgcc cdacacccca ggagcgaatt cacacaacdc bobobbbboo tgatggcggg gatggccggt cttcgcctgt tettggegee ccatccggat ctggaacac gactttttta tategaagae 266266262 ttegtatgee gtgaacgtcg ttaccggtgg catetacace actetgaaaa acaaccagac ggggatgcgc ageteatggg getgaatett cgccttcttt tctggggacg caccttcatc aacactagcc acacattgag aacaactacc tcaatgagcg 6606606606 441 561 501 141 201 261 321 381 481 541 601 661 721 781 841 901 961 021 081

FIG. 3A

cccggatatc accttccggc tggcgaaaga acgtgcgcaa tegaateega gttacgácga cgctgaatat ccggtaatgc caadaaaad ccttcgccgg aggacgactg acaccattac cgctgcccac gtgtgtatta ataaccaaaa geagaceat ctggtcgca ttcgcggcci acgatatoga aagggotgto cgcaqatcta tggcgctgga gaggtatgga cctccaqcat atgaoattat tgacatacaa cgacgetacg aatcccattg agagattaca ggctatctgg tcaatagccg cdcddcaaaa gccgatccgc gagatteagt gaaaccacta aatctegget ctctattttt agcagcatta aatcatgcga cagtatttac tatggcatct cacgicgate aggaccgggc atcaacaagg cttcgccagt caggatgett caaagatetg ggaaagecat aacactggcg agaccacaca tgaat taat t acagatcaat cateteetge adcccacaaa ccgtaccgta cagagtacca ttcaccaat ctattttacg atttgcccgc tgtgatgcgc gaatctgcat caaagaaggc agocacacac gacccgcgat ttacatccgg ttcttacatt ggacatgtat cgcgctataa tccaggacga ttcgcccta aacggtacga gcgtggtgat actttttca tegggeatga aagatacggt cggacattgg cagaaattt gccatatacg aggegt tgga caatgegget ccadcddcda tgggccagat tgcgcgacga ccggttcqat gcgtatcctg ttgagattcg tggaacaaga tcagggcgct atgaaaacct cctatgccga ataccaccac agtecetgta ctctctctcg atggcgctct gccattagcg aacgeeegtg atcatgtttg aacctgtgct tatacccaca atggattcac teggacatga gccataggta ggttcgccgg tegegetata acadcdaaad tggctaaagc gtggccaaaa tttatgacca attattgata gtgcatactt gtgccgccga ttttccccg aaaggtatta attgaagget cgccatcaac gatcaaaacg aaacdcdcaa tggcgatatc gacggcggtg aacctatatt atatecetae taatatgage taaccttgac cgctcacgtc atttgcgcag cacctctcat tattgcctac ctggcatgcc deaacedaaa acgagaaatg ttgcaggcg Goododoooo geteaccetg teeggaaaaa tacctagcga aggtactgaa ctegtattag tcatccgatt 1741 801 1861 2041 1921 1981 2101 2161 2221 2281 2341 2401 2461 2521 2581 2641 2701 2761

FIG. 3B

gattececa gegtgeette aaatatattg ggcgttaaag tcaaatcgcg ttcqcqttca ccadaadcaa atttctatca attaaattaq tatqcqqaaa gccttaataa ccttacgggc actgeegaee tataaqtatc ttcgcgctgg gcaatcotta gatattccgg acgggactta cattacatca ctgttctatt ggaagcggta cacgcccgct ggttcatct acctacacc agaagcgtta cgtgaatcc ttcctccca ttgcatggca gctatcgctt ttatattggc tgtgctacaa cgccaataaa ttccqqctca tattcacctg actcacqaac gaattttaa tagggtagaa aaaggtaact agagtettt gttaaagett caatagggag aatatteatg gttatcgaat gttaatggca ggttgatgcc tattttagct tegegtgttt aatcattcta agatggcaga agcatccgca cagtaacttc tggctgccgg aaaaagtgcc atgatttett cgagtgacat aaaatatete agcacctcgc aaaaaacddd geegaacage ageteaceat gccgcggtaa ttcacgatta teegetaeae acgaagactg ttatttcaag aacatcgcag gcgcgccgtc teagetttat agacgaaaga agcgtgaaga aggegeagat gcgtctttt ctgagtatgt aaataacagg atattiggag caaatactgg ctgtcgaaca acactcaata cacageegat aaageettet gttgttccaa ttccgccgg tcccatgcct acgetgtgee cttcagcgta aagattgcca tatttctcca gatgaagcgg teggeaateg gacaacgaaa gacgaaaacc gaaaccgaag atcgaaagag tctgtataaa cgacataagt tattttctcc tgaggegtta gtctcagatt gctgagcgcc agaggcagtg actaaagaaa attgccgatg acaaaaacta taacgacgtc gccgaatgcc gaaaacagtc acgttacatt cactatccag aaacccacca ggaactgtac aacatatect atttttgctg catgggaaat aatcattege agetgetega adaadcaada accttatat attcagggtt aaaagggact cttacagttc ccggctgggt acctgggtta ccgcgctctc atgtgatggg atgggaaata caaatattac attcaggtgg aagtgaagaa cgccgcatga ceggettetg ggtaacatat cctggcagac gegatgaace accggctgac tgattcgttt agatageget atttgttgat 3541 3661 3841 3901 3961 4021 1081 4141 1261 3181 3421 3481 3721 3781 3301 3361 3601

F1G. 3C

ggacaggtac gtgctggata tattigicat caigggatta iccggcicgg gtgcgcagga gagagagege gatgaactt cctgatatel ggcaatcaac gcatatgacc agagcgtcgc gcctgattga acccacccgc cgcctacccg cagacgetga gettegegag cacteatace tegeggegea attacgetea cccgcgcgct gaaggegaga cttctcaatc cagicatitg ttagcgggca gggattgaga gttgggcttg gccaaaatat teegeeeteg acgecagtet ggecattgaa cggtatggaa aatggtacgc gatggtcttc gcgtcaggtg tgaagcgttt cgttgatatt gcgtcagcgt tattaatgga gtagatecae tgattgacgg acaagattgc atacggcatt tggacgcctt ccggtgggat 4501 4681 4741 4801 4621 4561

FIG. 31

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tggctttctg gtcgaaaaag geettttetg ttcagtgtg tactgaagaa tttqctqqc tgcacagett ttattctcaa ctcaaaaqaa caaaaaaaaa atatctaqta tegagaaaaa ttaaaatga aattacacct ttatatqat ttcaccctat tatggettat ctatgatett ctccctctta ttttqactt aaaaactcgc tttttatggc agatgatttc cttatcgaga attaacact aacttgctta tagaacaaac ctataaaaac cgaatgtcga gaaatcgtaa caaaaggada gatcaaaaaa gcctgaattt agattcact ttctcacttc ttttcgaca getgetgttg ttgatttaaa tatattggcc aatgcaaaat acagacttgc cttttgataa atggactcgg gcadaatatc ccgctgtgat ttttacata ttgatgaaca taatcacaaa cttaatatga ateggeactg ctaaataaga ctttaaaaat attattatca gtctagagaa aattatatgo gaaatcaata gcaaagttgg tgtttctaaa cgagcettte agaagtttca agctgatgtt tttccctagc tttggattta cttgtcccct tatttttctt ttttttacg aatagacgtt getgeteetg tegtggaatt t taccactaa cgaaagtcac gactagatat tttactcc cgcgtcgttt ataaacattg atggtacgcc tctaaaaac tgcatttaac gggttttcga ttctgaatta tagagatgga ttctaaatc tgetttggea ttatgaatcc tatattgtgt tacagttta gatcaaggag caccaacca atctatattt tttgagttta ttgagtgaag qaattettat attccgtcc actggacaaa aatgataatt gtcaaggact taattgaaat gacgatgatt ttattattt tcaaaaatca aatagcacaa tetttaattt aacacgaaat 661 181 541 241 301 361 481 601

F/G. 4

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acttctggcc ctccgcaacg qtqqtcaaca tcatcaatac catagaacca gagatagaaa aacteteega egagaacta attegactgg tttaatgata coataacata cctqcgctca tggaagcgaa catcaacatg accdcaaaca ccgtctgaaa aggattcagc gccctcaacg gcaagtaagc accaataaaa aaaaaaccct ocacaaccac atcttacatt ctaaaqttt gcgaagtact gcatettiga talgcateae ttegaeatte aattaetegg eggtalggtt taagcactt tegeaatgge attttattaa gtegtataaa gactgcaggc tttaggtttg aagcdacaac tgattactgg caddcdcaac ccccacaadc atccgtgctg aacqatcqca ccctqcqccq qatqcqcaaa ategggatgg ttggccttgc ggaaggcacg agattattaa gctgttttct ggtagacget ctagaaaaag agtacataa ccagtcgttt cctttacttc ddcdcaaddc ggcataacga catttatact gccaggtgaa cggtgcgaac ttgcggcgag catetttett gegeattta tegtgetgtg dedecededa ctttggtcaa actattccgt tttgcctctt tgctgaccca tatgctaatc tttgaatatg tcatacacat gegaeagttt ctttcgccgt actacatege egggatgegt ctataaaaaa caacaataaa gagattttat ccacagaatt acteteteeg ccgaattcga ctgaagaatc ggataagcca ttatcttcat getaaataeg ttagggatgg ccaaagttaa ggtaatccgt ctcagcgcgc atccagaag tgacgcgctg cgaaccaaac cattgattat caaatctggc gcatctctta cacacaccac cagetatact tttataagag tecqeacaac gageetteag ccattegeae ctgcccgttg agggttatcg cctcacctaa cggggcgttt c.t taacqaac acaacaccac actggatacg acacccatct aaaabbbaaa agatttgtgc agtagaatac cagtagtegt ggaaaatcta 101 151 901 01 201 251 301 351 401 451 501 551 601 651 701 751 951 051

FIG. 5A

tetacatgae tgaageggaa aaaatteagg egateattga agatateaga acquactque camaqeeqqt attanqeaca aacaacaaca etacacate atacaetatt tacccataac atcaactaca tagaagcaga ccqqacctqq gaacgtactg cgaaaggeca geeggtgetg gtgggtaeta tetecatega qaccacaacc ctccctcctt acctgaacac actaaccaat aaagacatac cogcacetaa tecateaaga aaaagaagae teegaaaeet teeagggega tacaccacat tegitaggga iggigaggit aleategita acgaacaeae eggicalaee atcaqtatca acctaccaga catgecagea eeggeaaage gegaagetta egeagetgae ateaettaeg tacacaacaa cataacatte tagtagacga atcatttcca ctgaccgaac taaaagaga catcatagat tecagaacta ettecateta tatgaaaaac tageggggat gaeeggtaet cdadaacaac geceggeaga agacageteg gaaatgtata aaegegtgaa taaaattatt tegateacet atttagetea atetacaage tagataeegt cattatteca accaaceate caatgatteg taaagatetg gaagggagt ctctgtactc tccggccaac atcatgctga cactatacac caccaaacta agiggacice atecigateg algaagegeg tacacegeig ccagataaac agaaggtgtg cagatccaga acgaaaacca aacgetgget acqteetgaa egecaaatte eacgeeaacg aageggegat aacataacac catecactat ttaaatteet taacetaaet cagatacage tttaactace agccctgaag aacgtgtaca gcgtaaactg aggecactte teggtagaeg aggateteg atgatetgat getgattgaa gaactgetga atgeogggee gtegetggte egatggtetg tacctagage actgatacca aagetttega aaaatcagaa ctaatatcaa catcaacdac qtaccaacaa acqtaqttac 2251 901 951 451 651 701 751 801 2051 851 601

FIG. 5B

<u>cateacgata eggtaetaga ageaggtage etgeatatea tegataeega</u> <u>aggtatgaag ccaggcgaag ccattgaaca cccatgggtg actaaagcga</u> ctagatagag acagattate eggetaegat gactategeg accagtatag egggtegtag accacactaa <u>aagateegae egeagageaa attgaaagaa ttaaageega etggeaggta</u> cattetaate agaagataca ctaatacata tititaciic caaccaaqia iccaacataa iacataaaci cattcataag gagacattta tagatatoaa caatataaac gaaaccatta acagcatica tgaagatata ticaaaqcga ccatigaigc ciacaticca aggaacgtct geatgaagag acgetacata acageattet ageacagtee ategaaatat ateagegtaa agaagaagta attggtgetg agatgatgeg ctatagaga cctqcqtqqc atttacaaca atactagaat cattaaaata taaaattatc aatacactaa tetgeageeg cagetgeact gaeggegeag aceggagage cattetecat gatcagcatc tacgcacaga aagatccaaa acaggaatac aaacatgaat gtaacttcga tatctgcgtc agggtatcca acttaactcc categiatag aageegageg ittagegeaa aigeageage attacacaat ccqqqqctqc taccaaataa ggcagaagtt acctatcgat ttgaggaget tcgataacca cataactaac agtacgtata cctgaagaga attaaaaacc tactacaaac atagetagea teceatttet gtaggatatt ttegaceteg atttgecaat teceqteqta ccadcataaa aatatqatqa ctcccaqcat aacgaactat aggacatca agcacctage agcgatagac tagaagaaat atgeteggtg tactaattct acatcacqaa agataacaac ccacaqtcac gcaaagttca qtcaqqqqq caactactag aaccadaact teacttegag ttaccaacac tacaaatatt gaagaacgat 2451 3101 2701 2851 2901 3051 2801 2601 3001

F/G. 5C

gtgaacttca gaaaaactgg 3451 gcaaagtagg acgtaacgat ccttgcccgt gcggttctgg taaaaaatac aagtaaaagg aaaagctgca tttataacge tcccggcggt aagacaatga ctaactgttg caatgaaatc aaaattgaaa tgggtgaaac gccggaacag gcggtggtgc ttcgctattt gtegegeage agatgegeae atggegaata aactggagtt aattgeggta ggtattatte geaaegagaa 3501 aaqcaqtqcc atggccgcct gcaataaaag cgcaggattc tgcgcctttt ttataggttt gggattaccc cccaacattt ggaagaagtc aatatgaatt 3651 3551 3601 3701 3801 3751

FIG. 5D

agaaaccctc ttcgaccgac catctacctg cggtcgtcaa cactacggcg decadedeca acgacggtgc daacaccdd gctggttgga gegtgategg gegeettgge aggcgaaaa ccgcctggcg **oooboobbob** caagaccetg acggcgtgca gagtggatgg atgicggcac ttcaccacc tcacctacgg tctaaagact categatgat cacaaatco gatgcgagg tegeegeeag gggcttcagg tcggggtgat ccgcaaggtg gcacacaatc agtegeetta cgccgagctg aacdccdaca gaggacagat cttgaatcag ccgcggtggt tegeaceaaa accagatgga accdaacddc gateegtetg cgaagttgct gtggcggact agaaaaaccc 6606060006 tcgacgtgca ggtgatgggt ccggtgaagg ctggccggca acctggctaa acgcgacagt accaadacaa aactcaccga cctcaatgcg gagatgaaga ctgaacgtaa gtaaggatcg cctcaagaag gcttggagcg ggtggcctat gcgcgtgggc ctatgccgcg accadccdad gacgcgctct cdacaaddac actacggett ctaccatggg gctgtgctgt ctggccgacc cttcgccgtg ggcggatcgt agaactegte aactcgacca ggagatcacc tggtcaagcg gtcaacgact ccqcttcctc ggtttcgctg tettgtteta cggggacata gatgicgaga cageggeegt caacgttgcc tgcccgctta cegacagett gtgcgccgcg cacgcctacg caagcaacaa tcaacaccag gtcggtcgat cgggageetg cgctcgtcac tgcccgaggc atgaacgccg gatctacggc gccgcgtgca gagaacaact gaaggtegea atgacacccg ttcgacgtcg acadaaccc ctaccaccag ttgtccgac ggtgctggac gcacctaga categieaee gaggectgeg actggagagc obobobbbo cctacacatg gacgacctgt acctatatat gtcagcacat cgtcgtcgag sacgaettet ccgacgagtt ccaageegat 301 451 701 251 351 551 601 901 651 751 801 851

FIG. 64

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ctcaacaacg gtcagcatcc gatetacaag ccgagcgcta caatgigete tggcgggccg ddcaccdaca gctgcgcgaa caaggaagta acgagtegeg tgicgacgag acacacatc teggeatega categieege tgctgatcgg aaddadcacd gctgcagaac ccdcccadac gagaaatag tcatctccqq ttggcgccgc atggcgcact gcctggcact accccactaa acaaggacta accagccagg caccagagta gcatecegea agccagtcgc gtgctgggca ccgagcggcc agttcgccgg ggtcagctat ccaccatcac atgaccggca gctgggcgtg gacgacgtcg atcatcgcgg ccgatcagcg acaaddadda acacaacaac atctacacaa categaggee gtacgaggcg gaagaccagc agteegaeet attacgccat obooobbbbo cacagacacc ttcagccgcg cgacgagttc cagacgctgg getegeegge cgtgaagacc cgcggtggtc tgateggeae accaadcadc gactttctca cgcccgagga gaggaagcca tggtacaccg ggaattegte actegeegtt tegaccagge agatetacaa agaggcgacc tegactacet tacgaggtcg ccaccaacat gagtttgggt geggeagtte accacdadca cggcaa¢gtc tcctgatcga ggcctccaac daddccdcca caaagaactg aacdadddca agccgagaac tetacgacaa gccgatgatc ccaagtacat cagccagtac gtcaccgtcg ccggtggaga catcatcaaa geggetgtae tctggtgcag tgctcatcgt gagetgeacg ggacgtccac agaagggtgt cactggatga gtcgattcca ccgccgctac tegagateaa .acttccggc geoggagaga cacaaaagaa 2662662622 tgtgctggg cegaactgee ategaggeeg caccaataac sacctatac ctctgaagge gatggtgagg agaggegge cgaccaacat agtatetate cggcctggat poboobooq gatggaaaa gegtgeacg aacgccaagt 251 451 501 651 701 801 851 901 951 2001 2051 301 401 551 2101 601 751 351

FIG. 6B

aggacccgg cgccgcttca dedeceagae accedacaac ctcaaatacq ababbaabab tggtccgcga tatgccgaag ctatccggag tegagegega dacdccdaac tagaccgtaa ggtateggge gcatgagagc aatcggtcgg ccggttgccc agecgeage aagaactca ctttgaccta ccddcdcdad cgccgaaatc cggcgaggat aacggcggtg tetacaccga bboobsoobso cgagetgatg gccatcaaga caagaacgtc gegetggaea cctcaaggag ctcaacgtca tegagtacca tegeegeege ggtgcagcgc ccggtgctag ggctgaacct oddaadaadaa gaccacaaat actactcaaq aggaaatcgc atgaaagagg boooobooo acacacacaa gagtegeeeg ageceage tcaaaaccct gtggccggtc cgcaaggtca cgctgggtga ttattgacca ttgaggtccg dacccdcaad agatggacta gateegttgg ggtcacccgg caaggaccag acddcdcdac tggacggcac tgctggaggc gccgaactcg caacgtgctg getegaegge gtcgatggtg ggagtgccgg tggaggcggt cttgccgaat tattgccago geteggetea acaaddccdc aaccagt tac ttctatttgt cttggagacc daaccadcad cacctetacg ggacgcgttg agcacage acccacadad geggaggatg agedecagee aagccaagat cagcagaact cgcgaggagt cgcacgggaa agctggaacg acacaacaca dedecaaddd boobooobob gegaaaacet gcctacgtcg ccgactcgct aacgtcaccg tcatggccat cggctgccga geggategae ggagtcgcgc atggcgcggc gtgccgatcg ccaggtcgag atcctcgaag gggatcaccg gcgatgcgcc gtggcgtgaa gracetatae cttcctgttc acagacaaca acgaggtgat attgggatct cgateteace tgcgcgcgat agtgcattac tecggtece dadcccacaa tgtcatcacc tacgacatgt cggcgcgaac 2401 2601 2501 2801 2951 3001 651 2851 2901

-/G. 6C

cggtttctca aacggetata cgcatgcaca gttcgtcgac tttcctggcg tccagcgttc ggccgatttg caccattgcg tggcgttccg tegetegeae atcggcctgt ctggattctg atgggtgtat ttcccggata ccggcaggtt gagcagttgg agageggege bobboooobb aatctcaccc ggttcggcgc cgcgcacgct tgctggtgtt cgtgcctcgt ggageetgte taggttgcag atggttgcgg ggggttcgct ctgtccggtg **booboobbbo** cactagtict agcacaccc ctggtagcgg cggcagtgtc ctgcgatact caacategeg gacgacattc cgttagcgcg tacgacatcg cgccgccttc gggattagaa ggcagcgatc gegeegtege ggcatcgggg gtcacttccc gtcggttctg cgacaatcgt cggcgggcaa agtcaagaag cttcccqtcc gttcccagaa acatgetett ttcgttgca ctgggccgca ggtcgttga ggggtgcggc cttcgtcggc tetegaet teegggetgt ggtgctgcag 3451 3501 3901 3951 3551 3701 3751 3801 3851 3601 3651

FIG. 6D

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gaccctgacc tggatgggcc gaccaccata gegeacteae tetecgeeeg gtgccgttcg aacccddcca ggttggacac cggtggtcta tcatcggcgg taaaqactcc gccttggcga agcgaaaacc accctcgacq accaccetac cctacggcac gcgtgcacgt cgacgaaggt gtagaaatt cctgccggg aggeacaeg ateaeggaeg gaacggccgt tegeettate aagttgctgc ggccgactat dedadaeeed gatgggtacg cgacagtgag gccgacatca DOOOOOOOO accaaadaac ccgtctggcg ccgagctgag aaacccadaa gtgaaggcaa gccgccaacg gggtgattt cgacaacatg acaccattat agatggaget tggctaaacg gegetetace tgtgctgtcg accgtgcccc acgtgcaggt cttcaggtcg gategttege caaddacacc accatgggag tcaagaaggt ctcaccgacg ccdaccadaa atgtagaccg caatgeeetg ggcctataac actacctgcg aggcaccatt 0666666660 acggettgat cgaggeette cgaccgttcg aacgactacc ggtgcagcgc gggacatagc gtcaagcgcc cttcctcggg tttgggttcg gaccagaagg gcctggggcg gategatgae tattctacga tgtcgagaaa agcaggctgg cgttgccgag ccgcttacct aacgeegggt ctgatcgacg gectaegaet gtcgtcacct gacgagttca acctgttgcc agttaccgtc gacttettet acacatgeg gtccgacga getggaceaa tgtgttttac gegtgeaceg acaccegatg cgattccatc cadacacca tegageggga caccaatatc ocdccddcac oboobobobo aggtegeatg acctgggcga caataacgag agatgatet 301 351 401 451 551 501 651 701 251 501 801

FIG. 7/

acgeaeegte aaggagcacg gctgcagaac ttcaccgagt tegeceggtt ggegtgeege teggeatega ctcaacaacg categieege tgctgatcgg acaaggacta atctacgcaai ccaccatcac categaggee gaagaccagc ggtcagctat **G**GGCCGGG cgacgagttc actegeegtt ttcagccgcg cagacactag ggaattegte tgcaccaggc gctcgccggg tacgaggtcg ctccaactgg gagaccacca tctaggagaa caaagagetg agccaagaac ggacgtccac agaagggtgt aacdadddaa tgctcatcgt ენეეენენნნ caacctgtac ccgccgctac tegagateaa tacttccggc ggcgtgcacg ctctgaaggc gatggtgagg ggctggttt 101 251 301 201

ttaccagaag tatgacacca attttaatcg ctatttcata getaatgtta tgattctaca tagctgacat gtgatatcgc aaddaaaaac tcagtctatt atttattagg :taggtaaac aqtcaatqaa ataacttett tgagctaggt atagggtaat acttattaac ttaactgat atgccaacat gatatattai tgggatttti aacaataaat taaqtttaaa qcacttqtg† ttagtgatao dedaacdaaa acqttattac gtactaataa ggtggactca cagattatga tagatgataa gtgtattcaa aacadcdaca tattctqaaq ttataatata aattaaacag acataaaatt attcataaag gctgagttat acttaaacag taagacgaca taatagtatg tagtgactat atatttatat caaacagaat adacggcaat attacttaca actccaaaag ttctaacta agctaaagga ataataaaga t tagaagaaa gtagaacatt agaggtgttc tgaagaaatg catggtgaat tcattgatga aatatatta gaaacaatte atgattattt tggtattgca cataaaatat ggetetaaae gaatgaaaca gttaagttaa tgtcataaat gtcggattaa tacqaqataa cattttgcaa gaatgaagta cttgatggca agtaatcgct gtgttcaaag cdcacaadac taagtataat acagttttt qtaataaaac aaaaadcaaa aaattatggg acaggtgaag tagettgatt caaactagtg agtegaaaat attagctggt gttttttat atattigaat gegtattgag tgttagagaa gtaattigti tttgattact gcgtccatta aatgagtaaa tataaagttc gagatgaga acttaaatge gtgaagcata t t t t qcacaa tgtaggccaa gaagaaatte tgataatgtc gtatagtecg ttgctgataa catatgcact tacttatcaa atggatgtat aggtttgact gttcgataat aacagaatgt atcaaaaatt aagtataatt 351 401 551 751 301 451 251 501 601 651 701

FIG. 8A

taatgatteg tgaatcagca aaaaatcatq agggcaacca taggtgaagg tcaaggtgat tttcaaattt tgccgttact cgacatgaat atatgatgta gaagtattaa ttctcqqaa ttcaaaatga agaatgtaca agaagaattt aaaggtaaat acgaggcacg tacgccatta attatttctg gtgaagctga aaagtcaacg aacaddacda acadaacaad ataaacctat gtgcgcacgt gacaaaaagg caagagetgt tctgaatata attetagaeg caagatgaat gageegaeta aggtacagag ggcgttcaaa aaactgaaga attecgaeaa cattagecaa gttgttgaaa aacacaaggc attaaatgeg aaaatqttaa tgtacattta tattgatggc caggccgtcg agactatttc gatatcaaat acagetttac ttgaaaactt aaaatggtat teggggtaca ttacgtggtc tttatcatta tagcagtaat agaaaatgat ggtacageta atttaattta agttgagact ttacattcca agtaactcaa gtcatgatgt aaacdaaagc tcatatcaac actatatgat cgtacaatgc agcgaaggaa gcaggcgctg tgtttttgcg atgitcaaag aattgaatca gtcgcttcta gaacgittac atggcgtcta agtagaagat gtactattgc acatggctgg ttaggcggtt tgatgaccag atgttattag gataagtetg cgtggtatcc aagctottga gggtatgaca cacaagcaaa tacgatgada gtgeggataa agetgaaegt cgtgacgtag atttacagga ataacatgac tgaaattgtt attgccacta cg tagaggaa aaaggggata tttggttct actetacace acaaca taac aacatgaagc ctegtegtat tgattataaa caaaatgttg tacattacaa gatttacacc gtgctattag acttaaaaaa tcactttata ataaacttgc :tgatgcagt tgtcgatca atctaaaact agaaatatt 951 851 901 201 251 351 451 501 601 651 1701 751 801 301 401 551

FIG. 8E

ggatcaatta tagagaattc ttttatttta caaqcqttaa gtatcttaga aacdooadaa aaagtcaaaa ggtgaagcga agcacgtttc gttaaaggeg tccaattaaa gtacttagtt tgcaatgcta cagcagataa atcttcttac atttacttc cattacqtga caaaatatta tgctgaagat ctgtagtaca agttgaagat gagaataaac tatattaata tatcatgatg gacagetete aagttgtaga aaggtaaaga aataactcct aagttgttaa acgttgttat cattaatgac gcagcatatc tgagcgtatg caacaaaatc gaaaccaatc gtggtagtgg tegacacaat aattatctat tatcaattac tcatcgacta ggaggagtet caaaaacgtg tagaaggtaa taacttcqac gatgatatca aattatttga aatgatataa tgaotgagtt ttettatgea atttaaaat taagattgaa actgatcata gattgtccat aacaacgtga aacagagtt aagtgaaacc tatg tacaacqtag gtattacgta tcgtttgggc gaaggtcatg ttgtaaattc atggaaaata ttgtgttatg tgatgaagaa tatcaaccat gtgaaaaaac ggtttaggca cattacagag gaagaacaaa t tcact tacg ggtaaagaaa tegtaacgat tagccattgg aaatgtgaag cattcaacat aagaaggtga atttcgaag cgtcaaggta cgageetgaa aagaagatac atcaagttgg atactttaaa atacgatgaa ataqtattat agatatetta aatattgaac aggattgcc ttaaaaaatc gttctattga ctatcaaaat agctgaagat cacctatagt 2301 2351 2701 2751 2801 2851 2901 2501 2651 2401 2601 2451 2551

gttcagatca aattgttgac attagaaata tataagcaaa cacttatcca gcacatgata ttttaaata agatgatgaa ggattetga teatettea raaagacaga atattigaaa .gagaaagcc gtaatgtgag aggedeac acaaagtaat gaagttacaa ttccqcttaa acaaaacacd teccageaa ccaaacaqtt tgctggaatt gagaacagg tacacggggi tgtgccgaat gtgaatgcac atgtaaaag cagaadaaca gaagcatttg accttatcca t t t taacaaa aadcaadctg tgatgaagaa tttcagaaat cgatgattta ttgattgatt tatccatgtg ttgttgtaag acatgaacca atcatgataa tactgatttc ttaccacca ctaatatgtg tttcttttt gttcataatt tacatettga atcctatgag ttgaaaccaa gttcgtaaat aatggtgaca caattettae agattgcaag tattgaaaaa aaaaatacaa gatgtattaa gtaagataaa agaatgggtt acgeetaagt ttaatatgac acaagcatta getgeaagag atgattatga ctgtgagaag aatattacct gcacaaaatc aagaaadcda atttacttgg aagaaagaa agaatgtcaa acagetgttg acaattcago aatgaacagt gagaaatcaa acttcactaa gagaatgaaa attaaaaaat catataaatg aggagagaac gaagaaatgt attecaagaa cgccattcat ttacaaatgg tgattgagac tttagaaga aagaaacgta tccgccggaa tecgitetaa aaacgtgtat cttcgtgcag aactgatggt aaataagagg tgagcggcat gcaaatteta ctaacaaatt tgcattttct ttgattaat casattaage t tataccaa gacaataaga attgaaatca agaagcggta atgaccatga acatatggtt gggctcttgc cttattaaga gtccatgtca ctccgatgag taacaaaaa taggcagttt ctcattagaa aaacaaadc caagataaaa :gaaggagct tgggtggtat cttgaacgtt gacaagatca tgtcaatcgt cteeggaaae gaaaacttat tgacgtgaca 701 801 001 1151 751 851 901 951 101 151 201 301 401 451 601 651 251 351 501 551

aacgaigccg acttatttaa acgeettage gcaaattoto atcagtcgga tatttacgcg cttataatgo getteattte cgcgtacacc tatacacaag caaggtgctg ataaagctga gttgatatta accaatttac aagatgatta taattatqat caccaadcda aacaataact tataataaat tagccggtat atttataata gcaacgcgat tgaagataga ctgttgttga ttaggtactg gtatcgatga aagctaaaat gaacacad aggettegae tgaatacttg aagcgtgaag ttatgcgtcc tttgaaaaac atcgatgaag tccttggttt aacatctctt tgaaggactt atgaatctaa attecgtaae actatacatt ttgattgtcg ctgttcaacg aactattgaa aagttegatg accaattett catgaacgcg aaggtgcagt cacaatcgca aaattaggcg aaggtgttga gaatcacgcc ttacagtcaa ttatataatt aacadaacaa ctctatttta gaagaacgtg acttatatga ataatgaatt ctgaaaaatc ttaaaagcag attaacagat ttacgtgcta tggagaagta cttccgtatg gtcgattctc cagatteaga aggaagaaga agaaadacca ccaaaaaadac acgaaccatc tacatttcac cgctaaaaac agaacgtcat cattaactac gtgcatgtta aatggccgag acagettate tataqtacaa gaattattca atgaggtega tcaggggaag cactaaaatg aatcaqtaca aagttagata caatacagca tggttgtaga tccaaaacta atgccaggtc gctaaaacag acaaattcca gaaaacata agaaggggtt ttttcatcag aacaagtgaa atgictiaga gcaggicaaa ttattggtac taccgatatt gaagataaaa aaadadaada agcacgtggt ttgaacttga agatattacg ataacatggt actatcatta attgattatt casatgttt gaaaaaaaag gtagattaca acgtatgttc aggtegaaca tcacacatat ttgaggetaa cctgacttga ctateacat igacag t tac gtgcgtcatg gacaggtact tagagattga ctggtcgtgg ggccttgctg agatattatt catatataca 1401 1451 501 551 601 651 701 751 851 901 2201

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tgaagatgga taatctggaa catcaagata gatattgaac acateggaag aattgctgcg tgactatgcg gtgaagtcaa atttcgata agccaatata taagagagat ggtettttt tgaagtttta acattagatc attctattga ctatcaaaat caccaattga cgtgttgaag ttatcgattc ttctgaacgt aagaagacd gaaageegti tagatagett ggetgteage aaagtataaa aataatagag ggatcaatta aaagataatc aadcacaaaa atgtatcagc attttattac cacttegega gtcaatattg agtagatgat tggaagaaat t t dcacgaag tcgtgaggat tgaccacaga tacgtttcgg tgcacaaaaa tagaatacga cgtaataata gatgacteta gatacgetet tggaattat gcagcggtaa gatatgaaaa ggaccactgt tatatgetaa agatgitite aaggtaaaga caattatcac caaddacaac accagttatt tetttetaag aaagcttatg cgaacgtatg ttatggtgaa aatgaagaat tegatacaat caacaaaacc tacaatgatg ctgttgaatc. tcattacaat aaacgtatct gagttgatgg attaggtatg gacgacaagg gaattggaga aactattga agttgtatta aaaatatgga acagaccata atcttgaaat aaaagaatat tagaaccaca aattgggtta ttcatacggt tatecatgeg ttattatata attttgtgga atcaaaggta taggattgga tctgaattag tcaatgaatt ggtcgttctg gtatctcgag gtgaaatcat tgatgggccg cgatgcacgt attacaagat ccgtttatta tccatttacg geteaattga caatccaaca aattaaactg daadddcacc cagcaaatat gtgataaagc agatgateet gtaaatagta agaagatgaa tggaagatgg aaadaaaad gtgcaatcag cccgaccaat cagtatgggc tctatttatc ctgcaaaaaa atcaaaaatg gtaacaactt cataaacaac agaatcaagt tcagttgcgt 3551 3401 3501 3101 3151 3201 3301 3451 2501 2551 2601 2951 3001 3051 2651 2801 2901 2851 2751

cteccagate ctactgcaag gacceteaca gcggtcgctc caaggacgtg cctctcqqcc cgtagaggc catageegeg ttttttctcc gacgtgccaa ctgcatcttc attecacaaa tatacttgac tgctatccag caccataatc ctctgggtgc ccaaaaadac ggtgtacagc tteetttgge eetgatgaet tectecttet cggggctcta gcctgctcgc actacacaga acctcqtqtq gcggcagtca agetgttcae gcgcgggcta tegageggeg tegagggeat ccgcctttcc tattcgcgtt tctacgaget aggtgcccgc oddacdcddc tagatcactt oddacdacdc tgcactcgcg ttttcgtcgt ccgcgtcgtg gaccgagcca atggacggaa ggccttaaga gggaccgatt cdcadcacca bobbobbbob gcggtggcgc gtggactggc atgattetaa gtgcacacgg ccddcccdca ttcccctag taccactttc ccadadcacc tgccccgacc gacctgtccg atcgaagtgg acccacaacc agcgcggacc atgicaacga tgcgcgccgc caccccacg ccgcgaataa aaaatgttcc ctacctgcgc cgctgccctg ctataagaag cggcgtcttc ggagaccgag cdacdaadcc cgtcgagtac ctttgagcgc gctggagttt tagaaacgac booboobbob tttatttggt cccgcctcct caaccteggg gcaggagtcc ccddcdcaad gtacgtgctc gagcgcgatg cgagaccgag ttctgcgta tagigitacgo ccaccaacaa cggcgattgc tcatcagccg tatttggcac acacgaactt agggcaccc cgctcttcgt aatactttta deddedeeda tgctgctctt acccggcggt tggccgaaaa teggegggga adcacacc accactaget actatatega tgagcgcgag tegtgaacgt atgaacctcc accatacage gtggaggeta stgcctcctc cagicaatgi agegtegeaa gcgcggctca accaadaadac atccgcaagg cacctatage gccgatgacc accaacdacc gacaactacc aacgacctgt cataaataca atcctgcatt gccctcggcg gcgtggaaat gcagagtcgg tectegtttg 541 141 61 8 241 301 361 481 601 661 721 781 841 901 961 021 081

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26662226666 cctcacggag gacgcggctg gcgatccaag ggccgaticc ccggtgcgcc ggagacgetg gagcaatgac cggtggtccc cctgggggtt ccttgggggc attectated egacaeaegg attcatctca cacaaccacc ccagcgadtc accatgatca tegaeggaga egiggigege ggtgacaaac cccacatege tgactccgat ggcgcgcttt tctggacgtt egegtetgtt gaaatggeea geegeeeage egeateetet catcagcgat gtcgcccgtg obopboobob ocaccdaddc deceededda geceeeteg tcgggggaac acgacgeeet agtacttttg agegeetgtg ttaccacat ccttcgttgc acacggactc ctggtcttgc cggaggeegt aggagtggct booooobbob acgtgtactg cccgggtccg cgtcctaccg ggccacgtct gggagtatgt tattatctca tecgetegee gcctcctttg teceeegete cccdadaacd gccgacgtcc ggcggctgtt ctcatgctgg acattoggoa gtggagatgc cacdaccacc gtcgtggcat ggccccgtgt 6600066000 teggaceee gatccagtcg gacteggagg gccacgtacg boobbobboo atacataccc ggtgagccgg cdacaadacd ggaggggett agaactcacg caaacccaca ggetteteaa etaegegete gacccgggag ttccaactgc ccccccagg ccctctcgcc pobboboobb gtccggcggg gcagatagat occcdddacc odddaccdcc tacccagacg deegaedeea ggttggggga caccetgggt 6666066666 cgactcggat oboopboobo cdadacccdc tccaatgtgg aggaagacgc tgcggatccg ენნნნეენეე tggtgctttc tggatgtttg ccgtcccct tcaagccgct catctaccag cgtccgattc cctcggacgt atgactccct acccacacdc ccgtgaaccg ccaagcgtgt ეეენგენენე ctccccgctt aggacccggg gcagcgacgg cggtattcgg gtcaacgggt ctggtgcacc cctccggtcc tegageeege caddddaadd aatggcgtga agcaactttg teggegggga teacaegeet cgtcatcgg occacacccc gccgtggccc acggcctacc ctgggagatg cgcgaggaaa gacgactttg cgccgctctg gacgaccggt tcagcggtag ეეეეენენ<u>ე</u>ნ atcggagccg cccctcctc cccgtcgaag 261 421 541 721 781 841 901 961 141 601 661

FIG. 11/

ggtctgcaac gcacacacta cccgccatg cctcgcccgc agecgecage agagagaga ggccgtgctc cttcaqcqcc cgagaagaac tcacggggag gatacccatc cttcgtcatg categeegge tgggcggctc cacgetacaa cggcatgcag atttcaggac caacgcacta ccagctggtg gcgttttggc tttggcctgc gtgggaaatg occcaggcc teggegagea agtctgccga agcacgaagc acctggtaaa tegeegaett ocdddadccc 2662666660 gatccagtgg catttgactt tgaagaccag tegeeggett taccgicgac tcagtgccat cgcacaacaa ccgacgtgcg tgategaeag ccatgggaat acadcacacc tttacctaaa aggggtcgtg acctggacgg gegaeaatat gtcatggggt tatgaggagt accadcaacd agcatgicgc tccaggcaga gtgaacatca gaagetgggg etggatetgg ctgtcggcga aggettegeg tacacccgca caccagatcg tccagctgct ctggtggcgg ccgtggcaca deggeeaega tacgacaccc gcctccaage aacctgcggt tgtctgatcc ctggggcgag tttaacgact gcccagcgct ctgattcgcc cctctacgac ctactacacc gegggeeate ccgatgcatc cggcaacgac ggtgatgetg tggattttgg cctgacggaa ccattacgac cttccagatg ccttgactcg gtacctggag ccgggacacc gcacctcgag cgtccatccg gaccetgaag ccacategee cgtgcaggcg cagcaaaaa t t tcaagcgc tgtgcgcagt ccactacatc catgetgatg tgcaggcgtg. aggatagga tttccaccg ggacccgcaa ccctgttcga agetetacea cggtgaaccg gcaccgagat tgaatctggc agtgcacccg acategeega ggctgcaatc tggagtccat gcggcatgcg tegggetatg gggtcctcgc cagacctatt cctatggcat caacctacct aggegaeeet ccctcaaggt gegegtgegt gaagtggccc atcctggccc gtcgagcggg gggcactaca gtcacatgga boobobobo occacacccc qqcctgcaca aacgggggca cgtccgaccg gagttcgaga ctgggaagcg ctgaacctgg accacaaca gtcatgcccg ctctggatgc caggagetgg ctgaacaaac obopoobbbo cagatgaagg tcaaagacg tecaacetet t teaagt tet 2641 501 981 2041 561 741 801 2101 2221 2461 681 861 921 2161 2281 2341 2401 2521

FIG. 11E

ooboobooob agaaggegga egggaeeete aaaaacaddd cdacdacaac caccaacctg getaaaggaa cdacdccaad ccggcgattc ocacacccc gcatgtatcg cgccggccgc ggagatgeta gatgcccacc tgtgtgcgga teegegeeag tggacagtct cccacccct agegeggaet tetttggegg acacgetect agggcgagtg ccccctgtt ttgtcgcgct ctgcgccca ctggaggtga gagggettig ctggagccca tatgtcacgg cacgcatata aacadcdddd cttcaaccac tttaagcgca ctgatcgacc acadacccc ccgcggtacg aacagccagt ctgccatg getgtgaeeg ctctccctg catgaccetg ccttctggtc caaggcgacc ggacgtcagc cagcaaaaca gaagcacctc ccagaagttg ggacgccgg cggcctgcgc getecegtge gggcccgtcc tgagetgege ttgactacga atagccaatc ccgtcccaag ccctggtccg gcaaggttcg agegetttee tgatgaaaca cgcagatete tgacccggga cgtttagcgg tgccgcaggc attatctgca aagaccgcgt atgtactact actgtcgtcg adccadadca ttagcaagg tacgtcgacc ccagceteca gccgcctcgg ctggaacgca cagtagtccg tgcgttcgcg ttcactggg 3181 3301 3361 2941 3241 2881 3061 3001 3121

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attcatggac teageacaeg tcacatette acccacctc aacdacadda ggaaagtcad ttagcgacag tggcgcgcgg cgggcccgtc acccaaddac ttccatgggg teggggeege actcctcqqa tacaacccaa booobbbooo cgggctccgc obboooooo gggttaccac gattattagc gctacccagt tctcaaacqt tggttccaat ttcgtcgcga dadaaddacd gaaacccaca tgcaaatggg ctacgaaccc tetecateca cctccccct gtggcctttc ageaccagae ttagattgac cctqtcqaca accatcacct dacddadacd ctccacggga gaggactegg gacgactccg cacaccacca acddaccccd cgtaccgccc gtgttttgca gcctaccgca tcttcgatct cgcgcggtct ggcggaacct catatectee aaadccaacd cgccactctc ccggtgctgt tcctttcggt ccgttcatcg cgtccgagcc 660606660 cggcgaccac caccaacacc cggggagttc cccgcccct ეენენნენნნ gtetegatee ctcqcqqtcq ccctdccccc cctgggcgcc ttccgcggcc cactgtggga tatgataatc cgtggcgctc ctccqaaacg **b**o**b**oooob**b**o ccgactccga gccggggata catgigicic igaaaiggeg aggagatagg acttgacggt ttcctgtcgg ggccggtgcg ccgccctcgc cccccctgg gegateeee ccaactgcag cccagacgtc aggetgtece gegatgeeag cgtcgtccga cggagacget acaactccaa agagagaaga gaacccgg acadecadee gagatacata gaaccaccat cccctgactc accctatat tacttegggg tcctqttgcg gtgattcgtc cctgccgcat cccgaggtcg gtgettteea caatgegget gacctcgagg ggggatggcc tccgtgggga caaddacccc tgcgcccgtc tcagacggcc gacgatgaca cgcgcgtcgg ccggttctgg gccgctgttt gacteceeg ggctcgggtt cgtcgcagat geceacegat atgtgtttgg acgttacacc aacaggtagg ggagteatgg dacddacccd ategeaecee gactgccgg cacgggccc agccaaccac cgcgactaca scacqaqtac ggcgactgac ggacgtggcg acccdddaad cggcgtgatg occdaaccc cgaggatacg cattateatt acdccccddc cagetttatt odcadccddc catttacat 181 361 781 841 901 421 541 601 661 721 961 021 141 481 081

FIG. 12A

acctaacata actacataga gggcatcgg taccaaccet aggacgcggt **Gacccaca** accteggaac ენნნნეენე ggacctgga ggcccaggc agttetttt ctaaccagge tegagaaget gggacgccgt aggagagcaa cggtccccc acgggttcaa ttcacctqcq agcgggggct cctggtccct acggaggacg actttgggct tgaagggcg atcatgttta ggcgaggagt cccatccagg cggctggtta gadatgitca gccatgctga caggccacca gccagcatca adcacdcdcc gtgctcaqaa aaaaacgtca agegeeetet gaccttcccc gggattctgg cggtttctgg ctaatgatco acactegteg gcgtgccggg gccgcaacg tgcgcccgcg tccaaggagg ttcggcgggc ccaccaccgg aagccccttc agegetgega caacatette teegecaect egaeggegag cgaaacgatc 6000066060 gtcgtggtgg qtccacccc cgccatcctc occeddedec caacaaacad ggcacagcga cgttcgggcc cgactttcac ccccdcctc cctgtgcctg ceteaagege cccgcatcgc cgggttcctg ggtagaccc atggatgcgc cacccdaac ggcggtggcg cttctgtcgg gtatgcgacg tegeatectg gcaacgtgag tategetege tggggttcgg cctttgagga tggcggcgca tgctggagta teggeagege aagagtttta acdacdccad cccgcctgta ttcgcgaaca ggcgacaggg agatgegaeg agategtgee gctgatacct agaacgcgga atctgaggga tgatccacag cagatgtaca Gangagaga aagcgcctga gactecetgg aaggaggeee gacaccagca ctcgaggcca cccgcgctca ccacgaacct gegetegetg acgccctatc cggcggtccg acddaaadda tacgactgtc ctgaagtacg ategeeetgg tacacgtcca accatcacca caggegttea ctggaaccct cgggaggcct tacgaccacc ctcacgcccg cgccatcgtg gtacgagcac gtgcgtgtac sctegeegge cctgttcttc cccctagaa cdaccdcdad accctggtg gcagtcggcg gtccgtcttc catgegeeae ccaccacctc ccgcaactac sacceteegg gaaggteetg gttcgaccgg gegegtgeee cctgaactac caacgcatac catccatacc cctgaaccc actgtgcatg cttcgggctg 2641 2521 2401 2161 2221 2281 2341 2461 681 801 861 1921 1981 2041 2101 561

FIG. 12E

gcagcgtgaa ctgcagccga cgccccagtg tacacctcc geeggeteea acetetaeae acgccataca actgggagcg cctcggccca gcaaggtgac ggtctgtggc cggccttcga ttgatcacag cctccaccct actactgcaa tetgeacaag gtcccgcata cacgcgtcgg tacacacage ttcgcggggc agcgcacgtt ctggagaccg agagcatgat acacacaca gagetegae atgcagggcc ggccgctttc gccaagcaat cctagatacc cgggacctga atgetacgee aaggaactcg cggttcaaga acgeteceeg acggggatgt gacaacatcg geggtaette gaaccactag ctccgagggc tgcaacctgg booboopoop gcccctata gtcgctctcg atgeteegeg gcgctgtgcg ascetgites geceeacae getettgetg acaccaaccc aaacagccac tacatctacg acacgcaagg ggcggccatt atatcatgat agacagcacg gggcattggc acgegatgga egggetegag tcacagagaa ggcggacggg ceggitatgee ceaeeeeega gaggaggacg tetecaaget cagaggaata getttgeece cetgtteace cggcctgaag cgattttggc ggccgagttc gtaccgggcc cgagtgggag gecagiteat egegeteatg cccctccgg tgcagaccgc ggggtcaggc gaccagtaac caccaacaac cttegetece teageatect gatacgaggg acccdccca tegacetgtg tccccgcga cgctccgatc ccaaacactc ggcggacgtt tacaatccat atctggagtc cggccatgaa agcgcagcat gcggggtgtt catataageg ctaatggtta cacdacaacc cggctcctgg ctcgtccacg cacccatcct agccacttta tcccgccgtc ggccccgatt atgggcctgg atgctgctcg obooobboob ctgcgcaaca gagacgetge ccttgcctgg gaactgetga catcaaccac cgtcggggac tgcgtctccc gtcagccagg actetgtatg gegaceaaca taagcaacag ggagategte ggcgtgcgtg adcccdcddc tegecatgga gegteeette ctttcgaac 5006665065 ccaggccctg 2666622666 agtacccga tetggeeega gtgcctgaag cgccgaggtg gaacacddc odddacddc ccaatccatg agctcgtgtt gateteggae ggtccgcctt agttcgcaag ctgcgcgctg ctacgaccag 3241 3481 3541 3781 3001 3061 3301 3421 3601 1841 3901 3361 3661 3181

ccgctccttg ggcgatgatc ccagaattta tagatetteg ggaaggaacg aacaggtcgg caccagaaac gcggcgacgt dacaaacaca gatetgtgag ggactgttgg caagegeega aagttcatcc gegtacetae 66c6cacdadd obopoobopo ggatatcgag atcgggttca gcggccatcg cageceetat gaaaacctgg t t tcacaaca attegegtea dadcaddaa actactcatc cccacacctg tggagcgcgc tccagggtcc ccaccadoto ggatgacgtg cateateaac cctcgggggc ggggccctg gggtcgaaac tcttattatt oobboooboo gatcccggag caccaccatc cctcatcage gatccatatg catgiccacc ctggggaaag accgataaat ccgcggacga cctggtgacg acattettea etaetaegtg gctggtgctc acaacaacta atttattac tttagaagag gggaggcggt aacgcgtcca ggcgcgtggg ggggcgtcgt acdecadade 9999^t99c99 ctccgttccg cgagtcacga acatcatca gcaccatcaa aatgcgactc agtegaaega tectgagtee ccctcagcct cgaaataaag tatctqtqct cegeetegtt tgctgggcct cgcgcgtttc gggcagtett gtacttgacg gacageteta ccaaaadda ggtgatatga cagggcctcc cggctgtttc agcacctcgt ataacaaacg 6663666363 gtcttttttg gtcacctgcc geggategee gccagctttc gatgtctaac 800888888 ttcctgtcgg gaacagaagg cgcgtctaca tatgtggccc cgggtgcggg tactacatet gcagccagtc dadaaaadad gagatgcata gggatggggg ccttgtaaaa tgtccggggc gegatatgae gacagecteg tgattattac ეენენეენენ cctcctgcgg gegatteage cctgatagat tctgtttqcc gcgcgtgtac gacccgacg odccccdac cgagtgccgc cccttgtage cggcctcttc catccactcc gctggaggcg categaggge gtctccggtg cggtcggcga gggctgataa atcggatcgg cccdaaacca tctaccactt deaceaacaa ccagaacccc tegeggaggt geggeetete ccgtgcatac ocdaddcddc agaactacgt atteegeeee ccaacttett ggtctgggcg gcatcgggag gccaggacgt gcategaggt acdaccaddc aggtggactg tcatgatcct tecgatecea 5461 5401 4741 1801 1861 4981 5041 5101 5161 5281 5341 1921 1561 1621 1261 1441 501 1681 321 1381

FIG. 12L

gtaggtgtta tggggacctg cacgagctcg gtaactcatc accagcaaca ctgttgggcc cagcataggc ggcctgggat gggccgcggt ეემნენენე cgcccgcgat **bobboooboo** cgacggacgc ggaactgcgg agcegeggtg tteegegace egetgegegt gegegteteg aggeegaeeg acagegacg aacagegeca acactegege ageaegteet tagtcgtcct tcccgccggg cgggcgccgt gtettetteg gttggcgcgc agaggacgac ggtctgtgtg gtatctccga 6666666066 aggagagaga cggtccagtg tccacgagat ctgcag ttggggtgca agaatttegg ccggcgacat atgeteteca t tgaag tacc ggcggggtga tccgggggct cggctcgggg 5641 5701 5581 5761 5881 5821

tacatatgeg gcataatgtg attcacacg etttaaaaa tggggtctaa gaggaattgg ttggtgggag atatgaactc gtgtcatggc aaadaccaac gtatgtggac tacaqqacat gegaattate aatcgtctga gcctgcaatc aacaaccadc ttattcagaa gageetttt ggaacacttt ccagagtcgt tagagtccat gcggggagct tggggattgt ggtgcgataa getattecea cgcgccattt tacagaaatg ccatttttaa tattggctaa ggcggacgtc attatactat agggtaacag agcaacaata atagtgtett gactgiccac ggatattcca gagtatatca cacattgact tattcattag tatactatgg tatatttca tgcaaccgag tgttcacggg atctttactt ggagaagcg tacattgcgt geggeagaga tgggatagac gacaggaagc ggccattaac cgcagacatc ctaccatatg atttttggta ccgagatttg tggctggaca ggccacgcca cacagaaggt gtacgtcgat acaggtgtta cacgcaacac agacctgaaa catattggga cadaatgega tgctacgcac getagaeee acceatttg ctgcgacaac tttgagcacc caaagaatat gcagagtcga cgaacgcctt tggaacaggc catcqqaata ttgcaactac acactccacc ggtgtggatt aaaccactgt tetttacact ttatgeteta gcggtgaggt aagaggttgg ctatgaccat aaccetggca gtgctgtaat actttgacat cgtctcgaga tgtatttagc aaactgtgcg actatcaacg categeacet ctttggattt tgaataccat ttgtcattcc gctgttattt cagaggitia decadadaed cgcctggagc ataqttaqaa aggcactacc gaaattatgg getettatgg tttttgacc gatgatactg tetatggeea ctactagact gtttatttcg aagccgctgt cgaattaaag agcgatgtcg tatgaagaaa tttcttagac gtgttggtta atatccaacg gcacctcgcc tegtttgaaa decagaeaga ggcctttatc ggatagactc tacaattccg actatctta tctcctaaag aggagtgtgt cggaatgetg cccagacctg ggaatatgag cgcgtgcaac tgtacaccta agactgtggg gactetattt t gcgc taaag ggggcaattt taccctaaac aacaacgetg aatggctcgc tactgcgcta ttccgatatt gattgatett aaccetcaac 261 781 241 301 601 661 901 961 361

aggggttttc tcaggcccaa gettacaage tgctaattaa tgtttataac tccacgcata caaccaacga tgaacaccta tgatgacctc gtttgccttt tggatatggc gtaotctage gtgtgccata tcaatqcqat aaaaccaatc ttaaaqacag taacaaacca tgaactctat atttgccagt gcctcgccat agattttact atteteagt gaccagacta tccatqtctt ttgtagggca gtcccgatat ttccaatttc cgagagtttt accggttttt ctagagaaac aacggggtgt aggetgttat tttaaagggt aatgtaacac ggggaagtac aatcttttgg aadaaddcaa aatattacaa acaatttta ggcgttgaag ggettgtaca dadadcadcd gaatgegege gcgtttggtg atgctggacc gatacggacc aatagcagaa cadetacade tatgcaaccc agttgtggga atttgagtac ctgcaaaatc agcatgeega egaaaeeeaa tgcctaccca aaacaatac acaggitacc tactgctacc ccagagccaa cagaattatg ctegaaaact acgeeggaat tecaceteca cgcaactgcc atcccagaaa cacgittitg tttccacag ccactageta boboobboob ggctacatac cgaaatggtc gcatgtacta tagtetacae gtatttctat gtctagecet caactgttaa tagacaagca tcagaagtat ttaaaacagc cgtttgtgga teceegeete aattatggcc gcaagctggg taccetttga tatettegte ttagcaaagt geegegetae gcactgacat ggaatteagg ctagaagetg cttaaaacag tgataaatgc acattaccaa ctaaactcgc tctatgcaca gaaattatcc agcacatatc atgegeacea gacagggetg dacadaaaac ageggagaee acacacaac atgccaacag ctcagtaagt gegggaaat cgtagaette ctcccaaaat gccgctctgg gcaacgetet cggggggaac acagacetat getatetacg cacttattaa aaggacgttg tgtactgcta agacacaga gacgagttat gatgtgtgcc caacgcctgg tggtcaccc cadcatcaac gctgggactt tcctqtttac tttgcccagg tttaacttc atctccagag agagaccagc caagtacagt tatageetat catgtgtgct tgagcctgct aatctttgtt 2401 921 981 2041 2101 2161 2281 2341 2461 501 561 681 861 741 801

FIG. 13.